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SEVERN
TRENT

STL

STL North Canton
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ANALYTICAL REPORT

EM SCIENCE (OH)

Lot #: A4I290193

Angela Hurley

The Payne Firm, Inc.
11231 Cornell Park Drive
Cincinnati, OH 45242

SEVERN TRENT LABORATORIES, INC.


Roger K. Toth / Fox
Project Manager

October 7, 2004

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CASE NARRATIVE

CASE NARRATIVE

A4I290193

The following report contains the analytical results for three water samples and one quality control sample submitted to STL North Canton by The Payne Firm, Inc. from the EM Science (OH) Site. The samples were received September 29, 2004, according to documented sample acceptance procedures.

STL utilizes USEPA approved methods in all analytical work. The samples presented in this report were analyzed for the parameter(s) listed on the analytical methods summary page in accordance with the method(s) indicated. Preliminary results were provided to Angela Hurley and Kevin Kallini on October 01, 2004, and October 06, 2004. A summary of QC data for these analyses is included at the back of the report.

STL North Canton attests to the validity of the laboratory data generated by STL facilities reported herein. All analyses performed by STL facilities were done using established laboratory SOPs that incorporate QA/QC procedures described in the applicable methods. STL's operations groups have reviewed the data for compliance with the laboratory QA/QC plan, and data have been found to be compliant with laboratory protocols unless otherwise noted below.

The test results in this report meet all NELAP requirements for parameters for which accreditation is required or available. Any exceptions to NELAP requirements are noted in this report. Pursuant to NELAP, this report may not be reproduced, except in full, without the written approval of the laboratory.

If you have any questions, please call the Project Manager, Roger K. Toth, at 330-497-9396.

This report is sequentially paginated. The final page of the report is labeled as "END OF REPORT."

SUPPLEMENTAL QC INFORMATION

SAMPLE RECEIVING

The temperature of the cooler upon sample receipt was 3.4°C.

CASE NARRATIVE (continued)

GC/MS VOLATILES

The sample(s) that contained concentrations of target analyte(s) at a reportable level in the associated Method Blank(s) were flagged with "B". All target analytes in the Method Blank must be below the reporting limit (RL) or the associated sample(s) must be ND with the exception of common laboratory contaminants.

The sample(s) that contain results between the MDL and the RL were flagged with "J". There is a possibility of false positive or mis-identification at these quantitation levels. In analytical methods requiring confirmation of the analyte reported, confirmation was performed only down to the standard reporting limit (SRL). The acceptance criteria for QC samples may not be met at these quantitation levels.

The matrix spike/matrix spike duplicate(s) for batch(es) 4275213 had recoveries outside acceptance limits. However, since the associated method blank(s) and laboratory control sample(s) were in control, no corrective action was necessary.

QUALITY CONTROL ELEMENTS OF SW-846 METHODS

STL North Canton conducts a quality assurance/quality control (QA/QC) program designed to provide scientifically valid and legally defensible data. Toward this end, several types of quality control indicators are incorporated into the QA/QC program, which is described in detail in QA Policy, QA-003. These indicators are introduced into the sample testing process to provide a mechanism for the assessment of the analytical data.

OC BATCH

Environmental samples are taken through the testing process in groups called QUALITY CONTROL BATCHES (QC batches). A QC batch contains up to twenty environmental samples of a similar matrix (water, soil) that are processed using the same reagents and standards. STL North Canton requires that each environmental sample be associated with a QC batch.

Several quality control samples are included in each QC batch and are processed identically to the twenty environmental samples. These QC samples include a METHOD BLANK (MB), a LABORATORY CONTROL SAMPLE (LCS) and, where appropriate, a MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD) pair or a MATRIX SPIKE/SAMPLE DUPLICATE (MS/DU) pair. If there is insufficient sample to perform an MS/MSD or an MS/DU, then a LABORATORY CONTROL SAMPLE DUPLICATE (LCSD) is included in the QC batch.

LABORATORY CONTROL SAMPLE

The Laboratory Control Sample is a QC sample that is created by adding known concentrations of a full or partial set of target analytes to a matrix similar to that of the environmental samples in the QC batch. The LCS analyte recovery results are used to monitor the analytical process and provide evidence that the laboratory is performing the method within acceptable guidelines. All control analytes indicated by a bold type in the LCS must meet acceptance criteria. Failure to meet the established recovery guidelines requires the repreparation and reanalysis of all samples in the QC batch. The only exception is that if the LCS recoveries are biased high and the associated sample is ND (non-detected) for the parameter(s) of interest, the batch is acceptable.

At times, a Laboratory Control Sample Duplicate (LCSD) is also included in the QC batch. An LCSD is a QC sample that is created and handled identically to the LCS. Analyte recovery data from the LCSD is assessed in the same way as that of the LCS. The LCSD recoveries, together with the LCS recoveries, are used to determine the reproducibility (precision) of the analytical system. Precision data are expressed as relative percent differences (RPDs). If the RPD fails for an LCS/LCSD and yet the recoveries are within acceptance criteria, the batch is still acceptable.

METHOD BLANK

The Method Blank is a QC sample consisting of all the reagents used in analyzing the environmental samples contained in the QC batch. Method Blank results are used to determine if interference or contamination in the analytical system could lead to the reporting of false positive data or elevated analyte concentrations. All target analytes must be below the reporting limits (RL) or the associated sample(s) must be ND except under the following circumstances:

- Common organic contaminants may be present at concentrations up to 5 times the reporting limits. Common metals contaminants may be present at concentrations up to 2 times the reporting limit, or the reported blank concentration must be twenty fold less than the concentration reported in the associated environmental samples. (See common laboratory contaminants listed below.)

Volatile (GC or GC/MS)

Methylene chloride
Acetone
2-Butanone

Semivolatile (GC/MS)

Phthalate Esters

Metals

Copper
Iron
Zinc
Lead*

- *for analyses run on TJA Trace ICP, ICPMS or GFAA only*

QUALITY CONTROL ELEMENTS OF SW-846 METHODS **(Continued)**

- Organic blanks will be accepted if compounds detected in the blank are present in the associated samples at levels 10 times the blank level. Inorganic blanks will be accepted if elements detected in the blank are present in the associated samples at 20 times the blank level.
- Blanks will be accepted if the compounds/elements detected are not present in any of the associated environmental samples.

Failure to meet these Method Blank criteria requires the repreparation and reanalysis of all samples in the QC batch.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

A Matrix Spike and a Matrix Spike Duplicate are a pair of environmental samples to which known concentrations of a full or partial set of target analytes are added. The MS/MSD results are determined in the same manner as the results of the environmental sample used to prepare the MS/MSD. The analyte recoveries and the relative percent differences (RPDs) of the recoveries are calculated and used to evaluate the effect of the sample matrix on the analytical results. Due to the potential variability of the matrix of each sample, the MS/MSD results may not have an immediate bearing on any samples except the one spiked; therefore, the associated batch MS/MSD may not reflect the same compounds as the samples contained in the analytical report. When these MS/MSD results fail to meet acceptance criteria, the data is evaluated. If the LCS is within acceptance criteria, the batch is considered acceptable. The acceptance criteria do not apply to samples that are diluted for organics if the native sample amount is 4x the concentration of the spike.

For certain methods, a Matrix Spike/Sample Duplicate (MS/DU) may be included in the QC batch in place of the MS/MSD. For the parameters (i.e. pH, ignitability) where it is not possible to prepare a spiked sample, a Sample Duplicate may be included in the QC batch. However, a Sample Duplicate is less likely to provide usable precision statistics depending on the likelihood of finding concentrations below the standard reporting limit. When the Sample Duplicate result fails to meet acceptance criteria, the data is evaluated.

SURROGATE COMPOUNDS

In addition to these batch-related QC indicators, each organic environmental and QC sample is spiked with surrogate compounds. Surrogates are organic chemicals that behave similarly to the analytes of interest and that are rarely present in the environment. Surrogate recoveries are used to monitor the individual performance of a sample in the analytical system.

If surrogate recoveries are biased high in the LCS, LCSD, or the Method Blank, and the associated sample(s) are ND, the batch is acceptable. Otherwise, if the LCS, LCSD, or Method Blank surrogate(s) fail to meet recovery criteria, the entire sample batch is reprepped and reanalyzed. If the surrogate recoveries are outside criteria for environmental samples, the samples will be reprepped and reanalyzed unless there is objective evidence of matrix interference or if the sample dilution is greater than the threshold outlined in the associated method SOP.

For the GC/MS BNA methods, the surrogate criterion is that two of the three surrogates for each fraction must meet acceptance criteria. The third surrogate must have a recovery of ten percent or greater.

For the Pesticide, PCB, and PAH methods, the surrogate criterion is that one of two surrogate compounds must meet acceptance criteria.



STL North Canton Certifications and Approvals:

Alabama (#41170), California (#01144CA), Connecticut (#PH-0590), Florida (#E87225), Illinois (#100439), Kansas (#E10336), Massachusetts (#M-OH048), Maryland (#272), Minnesota (#39-999-348), New Jersey (#OH001), New York (#10975), Ohio (#6090), OhioVAP (#CL0024), Rhode Island (#237), South Carolina (#92007001, #92007002, #92007003), Tennessee (#02903), Utah (#QUAN9), Virginia (#00011), West Virginia (#210), Wisconsin (#999518190), NAVY, ARMY, USDA Soil Permit, ACIL Seal of Excellence – Participating Lab Status Award (#82)

***EXECUTIVE
SUMMARY***

EXECUTIVE SUMMARY - Detection Highlights

A4I290193

| <u>PARAMETER</u> | <u>RESULT</u> | <u>REPORTING LIMIT</u> | <u>UNITS</u> | <u>ANALYTICAL METHOD</u> |
|--|---------------|------------------------|--------------|--------------------------|
| VE540/21-26/092804 09/28/04 13:10 001 | | | | |
| Acetone | 5.5 J | 10 | ug/L | SW846 8260B |
| Benzene | 0.27 J | 1.0 | ug/L | SW846 8260B |
| 2-Butanone | 2.3 J | 10 | ug/L | SW846 8260B |
| Chloromethane | 0.43 J,B | 1.0 | ug/L | SW846 8260B |
| 1,2-Dichloroethane | 4.5 | 1.0 | ug/L | SW846 8260B |
| 1,4-Dioxane | 720 | 50 | ug/L | SW846 8260B |
| Toluene | 0.33 J | 1.0 | ug/L | SW846 8260B |
| VE540/35.5-40.5/092804 09/28/04 13:30 002 | | | | |
| Acetone | 6.8 J | 10 | ug/L | SW846 8260B |
| 2-Butanone | 1.7 J | 10 | ug/L | SW846 8260B |
| Chloroform | 0.18 J | 1.0 | ug/L | SW846 8260B |
| Chloromethane | 0.24 J,B | 1.0 | ug/L | SW846 8260B |
| 1,4-Dioxane | 20 J | 50 | ug/L | SW846 8260B |
| 1,1,1-Trichloroethane | 0.48 J | 1.0 | ug/L | SW846 8260B |
| Trichloroethene | 0.39 J | 1.0 | ug/L | SW846 8260B |
| VE541/26.5-31.5/092804 09/28/04 13:55 003 | | | | |
| Acetone | 5.3 J | 10 | ug/L | SW846 8260B |
| Benzene | 0.24 J | 1.0 | ug/L | SW846 8260B |
| 2-Butanone | 2.2 J | 10 | ug/L | SW846 8260B |
| Chloroform | 0.27 J | 1.0 | ug/L | SW846 8260B |
| Chloromethane | 0.29 J,B | 1.0 | ug/L | SW846 8260B |
| 1,2-Dichloroethane | 2.4 | 1.0 | ug/L | SW846 8260B |
| cis-1,2-Dichloroethene | 0.90 J | 1.0 | ug/L | SW846 8260B |
| 1,2-Dichloroethene (total) | 0.90 J | 2.0 | ug/L | SW846 8260B |
| 1,4-Dioxane | 18 J | 50 | ug/L | SW846 8260B |
| Trichloroethene | 0.38 J | 1.0 | ug/L | SW846 8260B |
| TRIP BLANK/092804 09/28/04 004 | | | | |
| Acetone | 0.95 J | 10 | ug/L | SW846 8260B |
| 2-Butanone | 0.48 J | 10 | ug/L | SW846 8260B |
| Chloromethane | 0.22 J,B | 1.0 | ug/L | SW846 8260B |
| Toluene | 0.19 J | 1.0 | ug/L | SW846 8260B |

METHOD SUMMARY

ANALYTICAL METHODS SUMMARY

A4I290193

| <u>PARAMETER</u> | <u>ANALYTICAL METHOD</u> |
|----------------------------|-------------------------------------|
| Volatile Organics by GC/MS | SW846 8260B |

References:

SW846 "Test Methods for Evaluating Solid Waste, Physical/Chemical Methods", Third Edition, November 1986 and its updates.

SAMPLE SUMMARY

SAMPLE SUMMARY

A4T290193

| WO # | SAMPLE# | CLIENT SAMPLE ID | SAMPLED DATE | SAMP TIME |
|-------|---------|------------------------|--------------|-----------|
| GRDVE | 001 | VE540/21-26/092804 | 09/28/04 | 13:10 |
| GRDX1 | 002 | VE540/35.5-40.5/092804 | 09/28/04 | 13:30 |
| GRDX6 | 003 | VE541/26.5-31.5/092804 | 09/28/04 | 13:55 |
| GRD0D | 004 | TRIP BLANK/092804 | 09/28/04 | |

NOTE (S) :

- The analytical results of the samples listed above are presented on the following pages.
- All calculations are performed before rounding to avoid round-off errors in calculated results.
- Results noted as "ND" were not detected at or above the stated limit.
- This report must not be reproduced, except in full, without the written approval of the laboratory.
- Results for the following parameters are never reported on a dry weight basis: color, corrosivity, density, flashpoint, ignitability, layers, odor, paint filter test, pH, porosity pressure, reactivity, redox potential, specific gravity, spot tests, solids, solubility, temperature, viscosity, and weight.

***SHIPPING
AND
RECEIVING DOCUMENTS***

Chain of Custody Record

**SEVERN
TRENT
SERVICES**

Severn Trent Laboratories, Inc.

STL-4124 (0901)

S 4
14

| | | | | | |
|---|--|---|--------------------------|---|--|
| Client Address The Payne Firm, Inc. 11231 Cornell Park Dr. | | Project Manager Kevin Kallini | | Date 9/28/04 | Chain of Custody Number 163676 |
| City Cinti. | | State OH | Zip Code 45242 | Telephone Number (Area Code)/Fax Number (513) 489-2255 / 489-2533 | Lab Number |
| Project Name and Location (State) EM Science (OH.) | | Site Contact A.L.H | | Analysis (Attach list if more space is needed) | |
| Contract/Purchase Order/Quote No. Trip BLANK /092804 | | Carrier/Waybill Number — | | Special Instructions/ Conditions of Receipt * Appendix IX | |
| Sample I.D. No. and Description (Containers for each sample may be combined on one line) | | Date | Time | Matrix | |
| | | | | Air | Aqueous |
| | | | | Sed. | Soil |
| | | | | Unpres. | H ₂ SO ₄ |
| | | | | HNO ₃ | HCl |
| | | | | NaOH | ZnAc/ NaOH # |
| | | | | # | X |
| VE540/21-26/092804 | | 9/28/04 | 1310 | X | 3 X |
| VE540/35.5 - 40.5/092804 | | 9/28/04 | 1330 | X | 3 X |
| VE541/26.5 - 31.5/092804 | | 9/28/04 | 1355 | X | 3 X |
| Trip BLANK /092804 | | 9/28/04 | — | X | 2 X |
| J. M. H. 9/28/04 | | | | | |
| * Results to Kevin Kallini | | | | | |
| * Appendix IX | | | | | |
| Possible Hazard Identification | | | | | |
| <input type="checkbox"/> Non-Hazard <input type="checkbox"/> Flammable <input type="checkbox"/> Skin Irritant <input type="checkbox"/> Poison A <input checked="" type="checkbox"/> Unknown <input type="checkbox"/> Return To Client <input checked="" type="checkbox"/> Disposal By Lab <input type="checkbox"/> Archive For _____ Months (<i>A fee may be assessed if samples are retained longer than 1 month</i>) | | | | | |
| Turn Around Time Required | | | | | |
| <input type="checkbox"/> 24 Hours <input checked="" type="checkbox"/> 48 Hours <input type="checkbox"/> 7 Days <input type="checkbox"/> 14 Days <input type="checkbox"/> 21 Days <input type="checkbox"/> Other _____ | | | | | |
| QC Requirements (Specify) | | | | | |
| 1. Relinquished By Jean M. Hawry M. Hawry 3. Relinquished By Comments | | | | | |
| 1. Received By Kevin Kallini K. Kallini 2. Received By John Dohr J. Dohr 3. Received By John Dohr J. Dohr | | | | | |
| Date 9/28/04 Time 1640 | | | | | |
| Date 9/28/04 Time 1640 | | | | | |
| Date 9/28/04 Time 1640 | | | | | |
| Date 9/28/04 Time 9:45 | | | | | |

Severn Trent Laboratories, Inc.
Sample Control Record

RR280
 Client: 5670
 Lot #: A41290193

Case Number/SDG:
 Storage Location: MS

| Laboratory Sample I.D. | Transferred By | Date | Entered | Removed | Reason | Date Returned |
|------------------------|----------------|---------|---------|---------|---------|---------------|
| GRDVE | BURNST | 9/29/04 | Yes | | Storage | |
| GRDX1 | BURNST | 9/29/04 | Yes | | Storage | |
| GRDX6 | BURNST | 9/29/04 | Yes | | Storage | |
| GRDOD | BURNST | 9/29/04 | Yes | | Storage | |

**STL Cooler Receipt Form/Narrative
North Canton Facility**

Lot Number: A41-290193

Client: Pagan Firm
Cooler Received on: 9/29/04

Project: EM Science
Opened on: 9/29/04

Quote#:

by: JL KERD
(Signature)

FedEx Client Drop Off UPS DHL FAS Other: _____
STL Cooler No# K033 Foam Box Client Cooler Other _____

1. Were custody seals on the outside of the cooler? Yes No Intact? Yes No NA
If YES, Quantity _____

Were the custody seals signed and dated?

Yes No NA

Yes No NA

Relinquished by client? Yes No

Yes No

Other: _____

2. Shipper's packing slip attached to this form?

3. Did custody papers accompany the samples? Yes No

4. Did you sign the custody papers in the appropriate place?

5. Packing material used: Bubble Wrap Foam None

6. Cooler temperature upon receipt 3.7 °C (see back of form for multiple coolers/temp)

IR ICE/H₂O Slurry

None

Yes No

Yes No

Yes No NA

Yes No

Yes No NA

Yes No

METHOD: Temp Vial Coolant & Sample Against Bottles

COOLANT: Wet Ice Blue Ice Dry Ice Water

7. Did all bottles arrive in good condition (Unbroken)?

Yes No

8. Could all bottle labels and/or tags be reconciled with the COC?

Yes No

9. Were samples at the correct pH? (record below/on back)

Yes No NA

10. Were correct bottles used for the tests indicated?

Yes No

11. Were air bubbles >6 mm in any VOA vials?

Yes No NA

12. Sufficient quantity received to perform indicated analyses?

Yes No

Contacted PM _____ Date: _____ by: _____ via Voice Mail Verbal Other

Concerning: _____

✓

I. CHAIN OF CUSTODY

The following discrepancies occurred:

| | |
|-------|-------|
| _____ | _____ |
| _____ | _____ |
| _____ | _____ |
| _____ | _____ |

2. SAMPLE CONDITION

| | |
|-----------------|---|
| Sample(s) _____ | were received after the recommended holding time had expired. |
| Sample(s) _____ | were received in a broken container. |

3. SAMPLE PRESERVATION

| | |
|-----------------|---|
| Sample(s) _____ | were further preserved in sample receiving to meet recommended pH level(s). Nitric Acid Lot #052804-HNO ₃ ; Sulfuric Acid Lot #011-504-H ₂ SO ₄ ; Sodium Hydroxide Lot # -082404-NaOH; Hydrochloric Acid Lot # 100902-HCl; Sodium Hydroxide and Zinc Acetate Lot # 071604-CH ₃ COO ₂ ZN/NaOH |
|-----------------|---|

| | |
|-----------------|---|
| Sample(s) _____ | were received with bubble > 6 mm in diameter (cc: PM) |
|-----------------|---|

4. Other (see below or back)

| | |
|-------|-------|
| _____ | _____ |
| _____ | _____ |
| _____ | _____ |
| _____ | _____ |

| Client ID | pH | Date | Initials |
|-----------|----|------|----------|
| | | | |
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STL Cooler Receipt Form/Narrative
North Canton Facility

| Client ID | pH | Date | Initials |
|-----------|------|--------|----------|
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| | | | |
| Cooler | Temp | Method | Coolant |
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Discrepancies Cont.



GCMS VOLATILE DATA

SEVERN
TRENT **STL**

QC SUMMARY DATA

SW846 8260B SURROGATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Lot #: A4I290193

Extraction: XXI25QK01

| | CLIENT ID. | SRG01 | SRG02 | SRG03 | SRG04 | TOT OUT |
|----|------------------------|-------|-------|-------|-------|---------|
| 01 | INTRA-LAB QC | 116 | 115 | 88 | 84 | 00 |
| 02 | VE540/21-26/092804 | 118 | 112 | 93 | 84 | 00 |
| 03 | VE540/35.5-40.5/092804 | 119 | 113 | 90 | 81 | 00 |
| 04 | VE541/26.5-31.5/092804 | 112 | 113 | 86 | 81 | 00 |
| 05 | TRIP BLANK/092804 | 118 | 111 | 87 | 80 | 00 |
| 06 | METHOD BLK. GRKVP1AA | 115 | 114 | 88 | 78 | 00 |
| 07 | LCS GRKVP1AC | 103 | 107 | 101 | 115 | 00 |
| 08 | LAB MS/MSD D | 105 | 106 | 97 | 113 | 00 |
| 09 | LCSD GRKVP1AD | 103 | 104 | 102 | 116 | 00 |
| 10 | LAB MS/MSD S | 106 | 106 | 101 | 116 | 00 |

SURROGATES

SRG01 = Dibromofluoromethane
 SRG02 = 1,2-Dichloroethane-d4
 SRG03 = Toluene-d8
 SRG04 = 4-Bromofluorobenzene

QC LIMITS

(73-122)
 (61-128)
 (76-110)
 (74-116)

- # Column to be used to flag recovery values
- * Values outside of required QC Limits
- D System monitoring Compound diluted out

FORM II

SW846 8260B CHECK SAMPLE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Lot #: A4J010000

WO #: GRKVP1AC
BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | SAMPLE CONCENT. (ug/L) | % REC | QC LIMITS REC | QUAL |
|----------------------------|---------------------------|-------------------------------|----------|---------------------|------|
| 1,2-Dichloroethane | 10 | 9.9 | 99 | 79- 136 | |
| Chloromethane | 10 | 4.4 | 44* | 48- 123 | a |
| Bromomethane | 10 | 5.9 | 59* | 64- 129 | a |
| Vinyl chloride | 10 | 5.2 | 52* | 61- 120 | a |
| Chloroethane | 10 | 7.5 | 75 | 66- 126 | |
| Methylene chloride | 10 | 10 | 100 | 78- 118 | |
| Acetone | 10 | 7.1 | 71 | 22- 200 | |
| Carbon disulfide | 10 | 8.0 | 80 | 73- 139 | |
| 1,1-Dichloroethene | 10 | 8.2 | 82 | 63- 130 | |
| 1,1-Dichloroethane | 10 | 9.3 | 93 | 86- 123 | |
| 1,2-Dichloroethene (total) | 20 | 18 | 89 | 82- 116 | |
| Chloroform | 10 | 9.6 | 96 | 84- 128 | |
| 2-Butanone | 10 | 8.0 | 80 | 28- 237 | |
| 1,1,1-Trichloroethane | 10 | 7.7 | 77* | 78- 140 | a |
| Carbon tetrachloride | 10 | 8.4 | 84 | 75- 149 | |
| Bromodichloromethane | 10 | 9.7 | 97 | 87- 130 | |
| 1,2-Dichloropropane | 10 | 9.8 | 98 | 82- 115 | |
| cis-1,3-Dichloropropene | 10 | 8.7 | 87 | 84- 130 | |
| Trichloroethene | 10 | 9.2 | 92 | 75- 122 | |
| Dibromochloromethane | 10 | 10 | 101 | 81- 138 | |
| 1,1,2-Trichloroethane | 10 | 10 | 101 | 83- 122 | |
| Benzene | 10 | 9.5 | 95 | 80- 116 | |
| trans-1,3-Dichloropropene | 10 | 8.3 | 83* | 84- 130 | a |
| Bromoform | 10 | 11 | 106 | 76- 150 | |
| 4-Methyl-2-pentanone | 10 | 11 | 111 | 78- 141 | |
| 2-Hexanone | 10 | 8.6 | 86 | 35- 200 | |
| Tetrachloroethene | 10 | 9.1 | 91 | 88- 113 | |
| 1,1,2,2-Tetrachloroethane | 10 | 12 | 116 | 85- 118 | |
| Toluene | 10 | 9.5 | 95 | 74- 119 | |
| Chlorobenzene | 10 | 10 | 100 | 76- 117 | |
| Ethylbenzene | 10 | 9.7 | 97 | 86- 116 | |

(Continued on next page)

SW846 8260B CHECK SAMPLE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Lot #: A4J010000

WO #: GRKVP1AC
BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | SAMPLE CONCENT. (ug/L) | % REC | QC LIMITS REC | QUAL |
|-----------------------------|---------------------------|-------------------------------|----------|---------------------|------|
| Styrene | 10 | 11 | 108 | 85- 117 | |
| Xylenes (total) | 30 | 31 | 104 | 87- 116 | |
| cis-1,2-Dichloroethene | 10 | 9.1 | 91 | 85- 113 | |
| trans-1,2-Dichloroethene | 10 | 8.8 | 88 | 79- 120 | |
| Dichlorodifluoromethane | 10 | 1.5 | 15* | 70- 130 | a |
| Trichlorofluoromethane | 10 | 5.8 | 58* | 70- 130 | a |
| 1,1,2-Trichloro-1,2,2-tri | 10 | 9.6 | 96 | 70- 130 | |
| Methyl acetate | 10 | 9.4 | 94 | 70- 130 | |
| Methyl tert-butyl ether (| 10 | 8.1 | 81 | 70- 130 | |
| Cyclohexane | 10 | 7.4 | 74 | 70- 130 | |
| Methylcyclohexane | 10 | 7.1 | 71 | 70- 130 | |
| 1,2-Dibromoethane | 10 | 10 | 101 | 70- 130 | |
| Isopropylbenzene | 10 | 10 | 103 | 70- 130 | |
| 1,3-Dichlorobenzene | 10 | 9.3 | 93 | 70- 130 | |
| 1,4-Dichlorobenzene | 10 | 10 | 104 | 70- 130 | |
| 1,2-Dichlorobenzene | 10 | 9.6 | 96 | 70- 130 | |
| 1,2-Dibromo-3-chloropropane | 10 | 7.6 | 76 | 70- 130 | |
| 1,2,4-Trichlorobenzene | 10 | 3.7 | 37* | 70- 130 | a |

NOTES (S) :

a Spiked analyte recovery is outside stated control limits.

* Values outside of QC limits

Spike Recovery: 8 out of 49 outside limits

COMMENTS:

SW846 8260B CHECK SAMPLE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Lot #: A4J010000

WO #: GRKVP1AD
BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | SAMPLE CONCENT. (ug/L) | % REC | QC LIMITS REC | QUAL |
|----------------------------|---------------------------|-------------------------------|----------|---------------------|------|
| Chloromethane | 10 | 4.4 | 44* | 48- 123 | a |
| Bromomethane | 10 | 5.9 | 59* | 64- 129 | a |
| Vinyl chloride | 10 | 5.0 | 50* | 61- 120 | a |
| Chloroethane | 10 | 7.1 | 71 | 66- 126 | |
| Methylene chloride | 10 | 10 | 102 | 78- 118 | |
| Acetone | 10 | 6.5 | 65 | 22- 200 | |
| Carbon disulfide | 10 | 7.8 | 78 | 73- 139 | |
| 1,1-Dichloroethene | 10 | 8.6 | 86 | 63- 130 | |
| 1,1-Dichloroethane | 10 | 9.3 | 93 | 86- 123 | |
| 1,2-Dichloroethene (total) | 20 | 18 | 92 | 82- 116 | |
| Chloroform | 10 | 9.5 | 95 | 84- 128 | |
| 1,2-Dichloroethane | 10 | 9.9 | 99 | 79- 136 | |
| 2-Butanone | 10 | 8.3 | 83 | 28- 237 | |
| 1,1,1-Trichloroethane | 10 | 7.7 | 77* | 78- 140 | a |
| Carbon tetrachloride | 10 | 8.6 | 86 | 75- 149 | |
| Bromodichloromethane | 10 | 9.7 | 97 | 87- 130 | |
| 1,2-Dichloropropane | 10 | 10 | 100 | 82- 115 | |
| cis-1,3-Dichloropropene | 10 | 8.9 | 89 | 84- 130 | |
| Trichloroethene | 10 | 9.1 | 91 | 75- 122 | |
| Dibromochloromethane | 10 | 10 | 102 | 81- 138 | |
| 1,1,2-Trichloroethane | 10 | 10 | 100 | 83- 122 | |
| Benzene | 10 | 9.6 | 96 | 80- 116 | |
| trans-1,3-Dichloropropene | 10 | 8.5 | 85 | 84- 130 | |
| Bromoform | 10 | 11 | 105 | 76- 150 | |
| 4-Methyl-2-pentanone | 10 | 11 | 111 | 78- 141 | |
| 2-Hexanone | 10 | 8.2 | 82 | 35- 200 | |
| Tetrachloroethene | 10 | 9.4 | 94 | 88- 113 | |
| 1,1,2,2-Tetrachloroethane | 10 | 11 | 115 | 85- 118 | |
| Toluene | 10 | 9.6 | 96 | 74- 119 | |
| Chlorobenzene | 10 | 10 | 102 | 76- 117 | |
| Ethylbenzene | 10 | 9.6 | 96 | 86- 116 | |

(Continued on next page)

SW846 8260B CHECK SAMPLE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN SDG No: 4I29193

Lot #: A4J010000 WO #: GRKVP1AD
BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | SAMPLE CONCENT. (ug/L) | % REC | QC LIMITS REC | QUAL |
|---------------------------|---------------------------|-------------------------------|----------|---------------------|------|
| Styrene | 10 | 11 | 111 | 85- 117 | |
| Xylenes (total) | 30 | 32 | 106 | 87- 116 | |
| cis-1,2-Dichloroethene | 10 | 9.2 | 92 | 85- 113 | |
| trans-1,2-Dichloroethene | 10 | 9.1 | 91 | 79- 120 | |
| Dichlorodifluoromethane | 10 | 1.3 | 13* | 70- 130 | a |
| Trichlorofluoromethane | 10 | 6.0 | 60* | 70- 130 | a |
| 1,1,2-Trichloro-1,2,2-tri | 10 | 9.5 | 95 | 70- 130 | |
| Methyl acetate | 10 | 9.4 | 94 | 70- 130 | |
| Methyl tert-butyl ether (| 10 | 7.8 | 78 | 70- 130 | |
| Cyclohexane | 10 | 7.5 | 75 | 70- 130 | |
| Methylcyclohexane | 10 | 7.0 | 70 | 70- 130 | |
| 1,2-Dibromoethane | 10 | 10 | 102 | 70- 130 | |
| Isopropylbenzene | 10 | 11 | 107 | 70- 130 | |
| 1,3-Dichlorobenzene | 10 | 9.5 | 95 | 70- 130 | |
| 1,4-Dichlorobenzene | 10 | 10 | 102 | 70- 130 | |
| 1,2-Dichlorobenzene | 10 | 9.4 | 94 | 70- 130 | |
| 1,2-Dibromo-3-chloropropa | 10 | 7.2 | 72 | 70- 130 | |
| 1,2,4-Trichlorobenzene | 10 | 3.1 | 31* | 70- 130 | a |

NOTES (S) :

* Spiked analyte recovery is outside stated control limits.

* Values outside of QC limits

Spike Recovery: 7 out of 49 outside limits

COMMENTS:

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Matrix Spike ID: LAB MS/MSD

Lot #: A4I240145

WO #: GQ2K61AC

BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | SAMPLE CONCENT. (ug/L) | MS CONCENT. (ug/L) | MS REC | LIMITS REC | LIMITS QUAL |
|----------------------------|---------------------------|-------------------------------|---------------------------|-----------|---------------|----------------|
| 1,1-Dichloroethene | 10 | ND | 9.8 | 98 | 62- 130 | |
| Chloromethane | 10 | 0.18 | 5.8 | 56 | 40- 137 | |
| Bromomethane | 10 | ND | 4.4 | 44* | 55- 145 | a |
| Vinyl chloride | 10 | ND | 5.1 | 51* | 88- 126 | a |
| Chloroethane | 10 | ND | 9.1 | 91 | 59- 142 | |
| Methylene chloride | 10 | ND | 9.8 | 98 | 82- 115 | |
| Acetone | 10 | ND | 6.7 | 67 | 45- 128 | |
| Carbon disulfide | 10 | ND | 11 | 107 | 69- 138 | |
| 1,1-Dichloroethane | 10 | 0.21 | 10 | 98 | 88- 127 | |
| 1,2-Dichloroethene (total) | 20 | ND | 19 | 97 | 86- 115 | |
| Chloroform | 10 | ND | 10 | 100 | 83- 141 | |
| 1,2-Dichloroethane | 10 | ND | 9.8 | 98 | 71- 160 | |
| 2-Butanone | 10 | ND | 7.6 | 76 | 71- 123 | |
| 1,1,1-Trichloroethane | 10 | ND | 8.8 | 88 | 71- 162 | |
| Carbon tetrachloride | 10 | ND | 10 | 101 | 63- 176 | |
| Bromodichloromethane | 10 | ND | 9.8 | 98 | 80- 146 | |
| 1,2-Dichloropropane | 10 | ND | 9.7 | 97 | 87- 114 | |
| cis-1,3-Dichloropropene | 10 | ND | 7.8 | 78* | 82- 130 | a |
| Trichloroethene | 10 | ND | 9.9 | 99 | 62- 130 | |
| Dibromochloromethane | 10 | ND | 9.7 | 97 | 71- 158 | |
| 1,1,2-Trichloroethane | 10 | ND | 10 | 105 | 86- 129 | |
| Benzene | 10 | 0.98 | 11 | 99 | 78- 118 | |
| trans-1,3-Dichloropropene | 10 | ND | 8.1 | 81 | 73- 147 | |
| Bromoform | 10 | ND | 10 | 102 | 58- 176 | |
| 4-Methyl-2-pentanone | 10 | ND | 11 | 107 | 82- 135 | |
| 2-Hexanone | 10 | ND | 9.0 | 90 | 81- 128 | |
| Tetrachloroethene | 10 | ND | 11 | 107 | 85- 121 | |
| 1,1,2,2-Tetrachloroethane | 10 | ND | 12 | 116 | 88- 116 | |

(Continued on next page)

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Matrix Spike ID: LAB MS/MSD

Lot #: A4I240145

WO #: GQ2K61AC

BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | SAMPLE CONCENT. (ug/L) | MS CONCENT. (ug/L) | MS % REC | LIMITS REC | QUAL |
|-----------------------------|---------------------------|-------------------------------|---------------------------|----------------|---------------|------|
| Toluene | 10 | ND | 10 | 100 | 70 - 119 | |
| Chlorobenzene | 10 | ND | 10 | 102 | 76 - 117 | |
| Ethylbenzene | 10 | ND | 10 | 102 | 86 - 132 | |
| Styrene | 10 | ND | 11 | 110 | 83 - 120 | |
| Xylenes (total) | 30 | ND | 34 | 111 | 89 - 121 | |
| cis-1,2-Dichloroethene | 10 | ND | 9.4 | 94 | 87 - 114 | |
| trans-1,2-Dichloroethene | 10 | ND | 9.9 | 99 | 85 - 116 | |
| Dichlorodifluoromethane | 10 | ND | 4.6 | 46* | 70 - 130 | a |
| Trichlorofluoromethane | 10 | ND | 7.4 | 74 | 70 - 130 | |
| 1,1,2-Trichloro-1,2,2-tri | 10 | ND | 13 | 131* | 70 - 130 | a |
| Methyl acetate | 10 | ND | 9.1 | 91 | 70 - 130 | |
| Methyl tert-butyl ether (| 10 | ND | 7.8 | 78 | 70 - 130 | |
| Cyclohexane | 10 | ND | 9.8 | 98 | 70 - 130 | |
| Methylcyclohexane | 10 | ND | 9.6 | 96 | 70 - 130 | |
| 1,2-Dibromoethane | 10 | ND | 9.8 | 98 | 70 - 130 | |
| Isopropylbenzene | 10 | ND | 12 | 115 | 70 - 130 | |
| 1,3-Dichlorobenzene | 10 | ND | 9.4 | 94 | 70 - 130 | |
| 1,4-Dichlorobenzene | 10 | ND | 10 | 103 | 70 - 130 | |
| 1,2-Dichlorobenzene | 10 | ND | 9.7 | 97 | 70 - 130 | |
| 1,2-Dibromo-3-chloropropane | 10 | ND | 8.0 | 80 | 70 - 130 | |
| 1,2,4-Trichlorobenzene | 10 | ND | 4.8 | 48* | 70 - 130 | a |

NOTES (S) :

a Spiked analyte recovery is outside stated control limits.

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits

RPD: 0 out of 0 outside limits

Spike Recovery: 6 out of 49 outside limits

COMMENTS:

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Matrix Spike ID: LAB MS/MSD

Lot #: A4I240145

WO #: GQ2K61AD

BATCH: 4275213

| COMPOUND | SPIKE ADDED (ug/L) | MSD CONCENT. (ug/L) | MSD % | % | QC LIMITS | | | QUAL |
|----------------------------|---------------------------|----------------------------|----------|-------|-----------|-----|-----|------|
| | REC | RPD | RPD | REC | | | | |
| 1,1-Dichloroethene | 10 | 10 | 101 | 2.6 | 20 | 62- | 130 | |
| Chloromethane | 10 | 5.8 | 57 | 1.6 | 39 | 40- | 137 | |
| Bromomethane | 10 | 5.8 | 58 | 26 | 30 | 55- | 145 | |
| Vinyl chloride | 10 | 5.4 | 54* | 5.7 | 30 | 88- | 126 | a |
| Chloroethane | 10 | 9.0 | 90 | 0.91 | 30 | 59- | 142 | |
| Methylene chloride | 10 | 9.2 | 92 | 6.2 | 30 | 82- | 115 | |
| Acetone | 10 | 6.6 | 66 | 1.3 | 30 | 45- | 128 | |
| Carbon disulfide | 10 | 10 | 100 | 6.2 | 41 | 69- | 138 | |
| 1,1-Dichloroethane | 10 | 9.8 | 96 | 1.9 | 30 | 88- | 127 | |
| 1,2-Dichloroethene (total) | 20 | 19 | 93 | 3.7 | 30 | 86- | 115 | |
| Chloroform | 10 | 9.8 | 98 | 1.1 | 30 | 83- | 141 | |
| 1,2-Dichloroethane | 10 | 9.6 | 96 | 2.6 | 30 | 71- | 160 | |
| 2-Butanone | 10 | 7.4 | 74 | 3.6 | 30 | 71- | 123 | |
| 1,1,1-Trichloroethane | 10 | 8.7 | 87 | 0.87 | 30 | 71- | 162 | |
| Carbon tetrachloride | 10 | 9.9 | 99 | 2.0 | 30 | 63- | 176 | |
| Bromodichloromethane | 10 | 9.3 | 93 | 5.0 | 30 | 80- | 146 | |
| 1,2-Dichloropropane | 10 | 9.8 | 98 | 0.96 | 30 | 87- | 114 | |
| cis-1,3-Dichloropropene | 10 | 7.8 | 78* | 0.010 | 30 | 82- | 130 | a |
| Trichloroethene | 10 | 9.9 | 99 | 0.80 | 20 | 62- | 130 | |
| Dibromochloromethane | 10 | 9.4 | 94 | 3.6 | 30 | 71- | 158 | |
| 1,1,2-Trichloroethane | 10 | 9.8 | 98 | 6.4 | 30 | 86- | 129 | |
| Benzene | 10 | 10 | 95 | 3.9 | 20 | 78- | 118 | |
| trans-1,3-Dichloropropene | 10 | 7.8 | 78 | 3.8 | 30 | 73- | 147 | |
| Bromoform | 10 | 9.8 | 98 | 4.6 | 30 | 58- | 176 | |
| 4-Methyl-2-pentanone | 10 | 10 | 105 | 2.3 | 30 | 82- | 135 | |
| 2-Hexanone | 10 | 8.3 | 83 | 7.7 | 30 | 81- | 128 | |
| Tetrachloroethene | 10 | 10 | 103 | 4.5 | 30 | 85- | 121 | |
| 1,1,2,2-Tetrachloroethane | 10 | 11 | 112 | 3.6 | 30 | 88- | 116 | |

(Continued on next page)

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I29193

Matrix Spike ID: LAB MS/MSD

Lot #: A4I240145

WO #: GQ2K61AD

BATCH: 4275213

| COMPOUND | SPIKE ADDED | MSD CONCENT. | MSD % | REC | QC LIMITS | | | QUAL |
|-----------------------------|----------------|-----------------|----------|-------|-----------|-----|-----|------|
| | | | % | | RPD | RPD | REC | |
| Toluene | 10 | 9.7 | 97 | 2.7 | 20 | 70- | 119 | |
| Chlorobenzene | 10 | 9.9 | 99 | 3.2 | 20 | 76- | 117 | |
| Ethylbenzene | 10 | 9.9 | 99 | 2.8 | 30 | 86- | 132 | |
| Styrene | 10 | 11 | 106 | 4.4 | 30 | 83- | 120 | |
| Xylenes (total) | 30 | 32 | 107 | 3.5 | 30 | 89- | 121 | |
| cis-1,2-Dichloroethene | 10 | 9.0 | 90 | 4.9 | 30 | 87- | 114 | |
| trans-1,2-Dichloroethene | 10 | 9.7 | 97 | 2.5 | 30 | 85- | 116 | |
| Dichlorodifluoromethane | 10 | 4.6 | 46* | 1.4 | 30 | 70- | 130 | a |
| Trichlorofluoromethane | 10 | 7.4 | 74 | 0.050 | 30 | 70- | 130 | |
| 1,1,2-Trichloro-1,2,2-tri | 10 | 12 | 124 | 5.6 | 30 | 70- | 130 | |
| Methyl acetate | 10 | 8.8 | 88 | 4.1 | 30 | 70- | 130 | |
| Methyl tert-butyl ether | 10 | 7.8 | 78 | 0.50 | 30 | 70- | 130 | |
| Cyclohexane | 10 | 9.7 | 97 | 1.6 | 30 | 70- | 130 | |
| Methylcyclohexane | 10 | 9.6 | 96 | 0.39 | 30 | 70- | 130 | |
| 1,2-Dibromoethane | 10 | 9.5 | 95 | 3.6 | 30 | 70- | 130 | |
| Isopropylbenzene | 10 | 11 | 112 | 2.6 | 30 | 70- | 130 | |
| 1,3-Dichlorobenzene | 10 | 9.2 | 92 | 2.8 | 30 | 70- | 130 | |
| 1,4-Dichlorobenzene | 10 | 10 | 101 | 2.0 | 30 | 70- | 130 | |
| 1,2-Dichlorobenzene | 10 | 9.2 | 92 | 5.1 | 30 | 70- | 130 | |
| 1,2-Dibromo-3-chloropropane | 10 | 7.3 | 73 | 9.2 | 30 | 70- | 130 | |
| 1,2,4-Trichlorobenzene | 10 | 4.5 | 45* | 6.3 | 30 | 70- | 130 | a |

NOTES (S) :

a Spiked analyte recovery is outside stated control limits.

Column to be used to flag recovery and RPD values with an asterisk
* Values outside of QC limitsRPD: 0 out of 49 outside limits
Spike Recovery: 4 out of 49 outside limits

COMMENTS:

SW846 8260B METHOD BLANK SUMMARY

BLANK WORKORDER NO.

Lab Name: Severn Trent Laboratories, Inc.

GRKVP1AA

Lab Code: STLCAN

SDG Number: 4I29193

Lab File ID: UXJ24281.

Lot Number: A4I290193

Date Analyzed: 10/01/04

Time Analyzed: 09:32

Matrix: WATER

Date Extracted: 10/01/04

GC Column: DB 624 ID: .18

Extraction Method: 5030B/8260B

Instrument ID: UX11

Level: (low/med) LOW

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, LCS, LCSD, MS , MSD:

| CLIENT ID. | SAMPLE WORK ORDER # | LAB FILE ID | DATE ANALYZED | TIME ANALYZED |
|----------------------------------|------------------------|------------------|------------------|------------------|
| 01 <u>INTRA-LAB QC</u> | <u>GQ2K61AA</u> | <u>UXJ24303.</u> | <u>10/01/04</u> | <u>18:12</u> |
| 02 <u>LAB MS/MSD</u> | <u>GQ2K61AC S</u> | <u>UXJ24299.</u> | <u>10/01/04</u> | <u>16:41</u> |
| 03 <u>LAB MS/MSD</u> | <u>GQ2K61AD D</u> | <u>UXJ24300.</u> | <u>10/01/04</u> | <u>17:03</u> |
| 04 <u>VE540/21-26/092804</u> | <u>GRDVE1AA</u> | <u>UXJ24286.</u> | <u>10/01/04</u> | <u>11:26</u> |
| 05 <u>VE540/35.5-40.5/092804</u> | <u>GRDXK11AA</u> | <u>UXJ24287.</u> | <u>10/01/04</u> | <u>11:49</u> |
| 06 <u>VE541/26.5-31.5/092804</u> | <u>GRDX61AA</u> | <u>UXJ24288.</u> | <u>10/01/04</u> | <u>12:28</u> |
| 07 <u>TRIP BLANK/092804</u> | <u>GRD0D1AA</u> | <u>UXJ24289.</u> | <u>10/01/04</u> | <u>12:51</u> |
| 08 <u>CHECK SAMPLE</u> | <u>GRKVP1AC C</u> | <u>UXJ24279.</u> | <u>10/01/04</u> | <u>08:47</u> |
| 09 <u>DUPLICATE CHECK</u> | <u>GRKVP1AD L</u> | <u>UXJ24280.</u> | <u>10/01/04</u> | <u>09:09</u> |
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COMMENTS:

VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.: SDG No.: 4I29193

Lab File ID: BFB207

BFB Injection Date: 08/16/04

Instrument ID: A3UX11

BFB Injection Time: 1309

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

| m/e | ION ABUNDANCE CRITERIA | % RELATIVE ABUNDANCE |
|-----|--|----------------------|
| 50 | 15.0 - 40.0% of mass 95 | 16.8 |
| 75 | 30.0 - 60.0% of mass 95 | 44.5 |
| 95 | Base Peak, 100% relative abundance | 100.0 |
| 96 | 5.0 - 9.0% of mass 95 | 6.9 |
| 173 | Less than 2.0% of mass 174 | 0.7 (0.8)1 |
| 174 | 50.0 - 100.0% of mass 95 | 81.3 |
| 175 | 5.0 - 9.0% of mass 174 | 6.1 (7.5)1 |
| 176 | Greater than 95.0%, but less than 101.0% of mass 174 | 78.9 (97.1)1 |
| 177 | 5.0 - 9.0% of mass 176 | 5.0 (6.4)2 |

1-Value is % of mass 174

2-Value is % of mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

| EPA SAMPLE NO. | LAB SAMPLE ID | LAB FILE ID | DATE ANALYZED | TIME ANALYZED |
|-------------------|------------------|----------------|------------------|------------------|
| 01 VSTD040 | 200NG-A9IC | UXJ23209 | 08/16/04 | 1618 |
| 02 VSTD020 | 100NG-A9IC | UXJ23210 | 08/16/04 | 1640 |
| 03 VSTD010 | 50NG-A9IC | UXJ23211 | 08/16/04 | 1703 |
| 04 VSTD005 | 25NG-A9IC | UXJ23212 | 08/16/04 | 1726 |
| 05 VSTD002 | 10NG-A9IC | UXJ23213 | 08/16/04 | 1748 |
| 06 VSTD001 | 5NG-A9IC | UXJ23214 | 08/16/04 | 1811 |
| 07 | | | | |
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VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.: SDG No.: 4I29193

Lab File ID: BFB232

BFB Injection Date: 09/14/04

Instrument ID: A3UX11

BFB Injection Time: 1321

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

| m/e | ION ABUNDANCE CRITERIA | % RELATIVE ABUNDANCE |
|-----|--|----------------------|
| 50 | 15.0 - 40.0% of mass 95 | 19.8 |
| 75 | 30.0 - 60.0% of mass 95 | 51.6 |
| 95 | Base Peak, 100% relative abundance | 100.0 |
| 96 | 5.0 - 9.0% of mass 95 | 6.6 |
| 173 | Less than 2.0% of mass 174 | 0.2 (0.3)1 |
| 174 | 50.0 - 100.0% of mass 95 | 74.6 |
| 175 | 5.0 - 9.0% of mass 174 | 5.6 (7.5)1 |
| 176 | Greater than 95.0%, but less than 101.0% of mass 174 | 74.5 (99.9)1 |
| 177 | 5.0 - 9.0% of mass 176 | 5.2 (6.9)2 |

1-Value is % of mass 174

2-Value is % of mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

| EPA SAMPLE NO. | LAB SAMPLE ID | LAB FILE ID | DATE ANALYZED | TIME ANALYZED |
|-------------------|------------------|----------------|------------------|------------------|
| 01 VSTD040 | 200NG-IC | UXJ23870 | 09/14/04 | 1348 |
| 02 VSTD020 | 100NG-IC | UXJ23871 | 09/14/04 | 1410 |
| 03 VSTD010 | 50NG-IC | UXJ23872 | 09/14/04 | 1433 |
| 04 VSTD005 | 25NG-IC | UXJ23873 | 09/14/04 | 1457 |
| 05 VSTD002 | 10NG-IC | UXJ23874 | 09/14/04 | 1519 |
| 06 VSTD001 | 5NG-IC | UXJ23875 | 09/14/04 | 1541 |
| 07 | | | | |
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page 1 of 1

FORM V VOA

1/87 Rev.

VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.: SDG No.: 4I29193

Lab File ID: BFB246

BFB Injection Date: 10/01/04

Instrument ID: A3UX11

BFB Injection Time: 0730

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

| m/e | ION ABUNDANCE CRITERIA | % RELATIVE ABUNDANCE |
|-----|--|----------------------|
| 50 | 15.0 - 40.0% of mass 95 | 18.1 |
| 75 | 30.0 - 60.0% of mass 95 | 49.6 |
| 95 | Base Peak, 100% relative abundance | 100.0 |
| 96 | 5.0 - 9.0% of mass 95 | 6.6 |
| 173 | Less than 2.0% of mass 174 | 0.4 (0.6)1 |
| 174 | 50.0 - 100.0% of mass 95 | 78.4 |
| 175 | 5.0 - 9.0% of mass 174 | 5.7 (7.3)1 |
| 176 | Greater than 95.0%, but less than 101.0% of mass 174 | 75.0 (95.6)1 |
| 177 | 5.0 - 9.0% of mass 176 | 5.1 (6.8)2 |

1-Value is % of mass 174

2-Value is % of mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

| EPA SAMPLE NO. | LAB SAMPLE ID | LAB FILE ID | DATE ANALYZED | TIME ANALYZED |
|-------------------|------------------|----------------|------------------|------------------|
| 01 VSTD010 | 50NG-CC | UXJ24277 | 10/01/04 | 0801 |
| 02 VSTD010 | 50NG-A9CC | UXJ24278 | 10/01/04 | 0824 |
| 03 GRKVPCHK | GRKVP1AC | UXJ24279 | 10/01/04 | 0847 |
| 04 GRKVPCKDUP | GRKVP1AD | UXJ24280 | 10/01/04 | 0909 |
| 05 GRKVPBLK | GRKVP1AA | UXJ24281 | 10/01/04 | 0932 |
| 06 VE540/21-26/ | GRDVE1AA | UXJ24286 | 10/01/04 | 1126 |
| 07 VE540/35.5-4 | GRDX11AA | UXJ24287 | 10/01/04 | 1149 |
| 08 VE541/26.5-3 | GRDX61AA | UXJ24288 | 10/01/04 | 1228 |
| 09 TRIP BLANK/0 | GRD0D1AA | UXJ24289 | 10/01/04 | 1251 |
| 10 | | | | |
| 11 | | | | |
| 12 | | | | |
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VOLATILE INTERNAL STANDARD AREA SUMMARY

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.: SDG No.: 4I29193

Lab File ID (Standard): UXJ24277

Date Analyzed: 10/01/04

Instrument ID: A3UX11

Time Analyzed: 0801

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

| | IS1 (CBZ) AREA # | RT | IS2 AREA # | RT | IS3 (DCB) AREA # | RT |
|-------------------|---------------------|------|---------------|------|---------------------|-------|
| 12 HOUR STD | 1659318 | 7.68 | 2028310 | 5.04 | 1038318 | 9.90 |
| UPPER LIMIT | 3318636 | 8.18 | 4056620 | 5.54 | 2076636 | 10.40 |
| LOWER LIMIT | 829659 | 7.18 | 1014155 | 4.54 | 519159 | 9.40 |
| EPA SAMPLE NO. | | | | | | |
| 01 GRKVPCHK | 1648603 | 7.68 | 2028404 | 5.04 | 862434 | 9.90 |
| 02 GRKVPCKDUP | 1659474 | 7.68 | 2044371 | 5.04 | 878061 | 9.90 |
| 03 GRKVPBLK | 1616266 | 7.68 | 1815827 | 5.04 | 657391 | 9.90 |
| 04 VE540/21-26/ | 1562768 | 7.68 | 1775894 | 5.04 | 690306 | 9.90 |
| 05 VE540/35.5-4 | 1533746 | 7.68 | 1755793 | 5.04 | 651571 | 9.90 |
| 06 VE541/26.5-3 | 1560887 | 7.68 | 1815392 | 5.04 | 683725 | 9.90 |
| 07 TRIP BLANK/0 | 1579425 | 7.68 | 1774228 | 5.04 | 650817 | 9.90 |
| 08 | | | | | | |
| 09 | | | | | | |
| 10 | | | | | | |
| 11 | | | | | | |
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| 17 | | | | | | |
| 18 | | | | | | |
| 19 | | | | | | |
| 20 | | | | | | |
| 21 | | | | | | |
| 22 | | | | | | |

IS1 (CBZ) = Chlorobenzene-d5

UPPER LIMIT = +100%

IS2 = Fluorobenzene

of internal standard area.

IS3 (DCB) = 1,4-Dichlorobenzene-d4

LOWER LIMIT = - 50%

of internal standard area.

Column used to flag internal standard area values with an asterisk.



SAMPLE DATA

PAYNE FIRM INC.

Client Sample ID: VE540/21-26/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-001 Work Order #....: GRDVE1AA Matrix.....: WG
 Date Sampled....: 09/28/04 13:10 Date Received...: 09/29/04
 Prep Date.....: 10/01/04 Analysis Date...: 10/01/04
 Prep Batch #....: 4275213
 Dilution Factor: 1 Initial Wgt/Vol: 5 mL Final Wgt/Vol.: 5 mL
 Method.....: SW846 8260B

| PARAMETER | RESULT | REPORTING | |
|------------------------------------|----------|-----------|-------|
| | | LIMIT | UNITS |
| Acetone | 5.5 J | 10 | ug/L |
| Acetonitrile | ND | 20 | ug/L |
| Acrolein | ND | 20 | ug/L |
| Acrylonitrile | ND | 20 | ug/L |
| Benzene | 0.27 J | 1.0 | ug/L |
| Bromodichloromethane | ND | 1.0 | ug/L |
| Bromoform | ND | 1.0 | ug/L |
| Bromomethane | ND | 1.0 | ug/L |
| 2-Butanone | 2.3 J | 10 | ug/L |
| Carbon disulfide | ND | 1.0 | ug/L |
| Carbon tetrachloride | ND | 1.0 | ug/L |
| Chlorobenzene | ND | 1.0 | ug/L |
| Chloroprene | ND | 2.0 | ug/L |
| Dibromochloromethane | ND | 1.0 | ug/L |
| Chloroethane | ND | 1.0 | ug/L |
| Chloroform | ND | 1.0 | ug/L |
| Chloromethane | 0.43 J,B | 1.0 | ug/L |
| 3-Chloropropene | ND | 2.0 | ug/L |
| 1,2-Dibromo-3-chloropropane (DBCP) | ND | 2.0 | ug/L |
| 1,2-Dibromoethane | ND | 1.0 | ug/L |
| Dibromomethane | ND | 1.0 | ug/L |
| trans-1,4-Dichloro-2-butene | ND | 1.0 | ug/L |
| 1,1-Dichloroethane | ND | 1.0 | ug/L |
| 1,2-Dichloroethane | 4.5 | 1.0 | ug/L |
| cis-1,2-Dichloroethene | ND | 1.0 | ug/L |
| trans-1,2-Dichloroethene | ND | 1.0 | ug/L |
| 1,1-Dichloroethene | ND | 1.0 | ug/L |
| 1,2-Dichloroethene (total) | ND | 2.0 | ug/L |
| Dichlorofluoromethane | ND | 2.0 | ug/L |
| 1,2-Dichloropropane | ND | 1.0 | ug/L |
| cis-1,3-Dichloropropene | ND | 1.0 | ug/L |
| trans-1,3-Dichloropropene | ND | 1.0 | ug/L |
| 1,4-Dioxane | 720 | 50 | ug/L |
| Ethylbenzene | ND | 1.0 | ug/L |
| Ethyl methacrylate | ND | 1.0 | ug/L |

(Continued on next page)

PAYNE FIRM INC.

Client Sample ID: VE540/21-26/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-001 Work Order #....: GRDVE1AA Matrix.....: WG

| <u>PARAMETER</u> | <u>RESULT</u> | <u>REPORTING</u> | |
|---------------------------|---------------|------------------|--------------|
| | | <u>LIMIT</u> | <u>UNITS</u> |
| 2-Hexanone | ND | 10 | ug/L |
| Iodomethane | ND | 1.0 | ug/L |
| Isobutanol | ND | 50 | ug/L |
| Methacrylonitrile | ND | 2.0 | ug/L |
| Methylene chloride | ND | 1.0 | ug/L |
| Methyl methacrylate | ND | 2.0 | ug/L |
| 4-Methyl-2-pentanone | ND | 10 | ug/L |
| Propionitrile | ND | 4.0 | ug/L |
| Styrene | ND | 1.0 | ug/L |
| 1,1,1,2-Tetrachloroethane | ND | 1.0 | ug/L |
| 1,1,2,2-Tetrachloroethane | ND | 1.0 | ug/L |
| Tetrachloroethene | ND | 1.0 | ug/L |
| Toluene | 0.33 J | 1.0 | ug/L |
| 1,1,1-Trichloroethane | ND | 1.0 | ug/L |
| 1,1,2-Trichloroethane | ND | 1.0 | ug/L |
| Trichloroethene | ND | 1.0 | ug/L |
| Trichlorofluoromethane | ND | 1.0 | ug/L |
| 1,2,3-Trichloropropane | ND | 1.0 | ug/L |
| Vinyl acetate | ND | 2.0 | ug/L |
| Vinyl chloride | ND | 1.0 | ug/L |
| Xylenes (total) | ND | 2.0 | ug/L |

| <u>SURROGATE</u> | <u>PERCENT RECOVERY</u> | <u>RECOVERY</u> |
|-----------------------|-----------------------------|-----------------|
| | | <u>LIMITS</u> |
| Dibromofluoromethane | 118 | (73 - 122) |
| 1,2-Dichloroethane-d4 | 112 | (61 - 128) |
| Toluene-d8 | 93 | (76 - 110) |
| 4-Bromofluorobenzene | 84 | (74 - 116) |

NOTE(S) :

J Estimated result. Result is less than RL.

B Method blank contamination. The associated method blank contains the target analyte at a reportable level.

Data File: \\pcapn04\\dd\\chem\\NSV\\a30x11.i\\J41001A.b\\JXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VES40/21-26/092804

Sample Info: GRINELIA,SHL/SHL

Purge Volume: 5.0

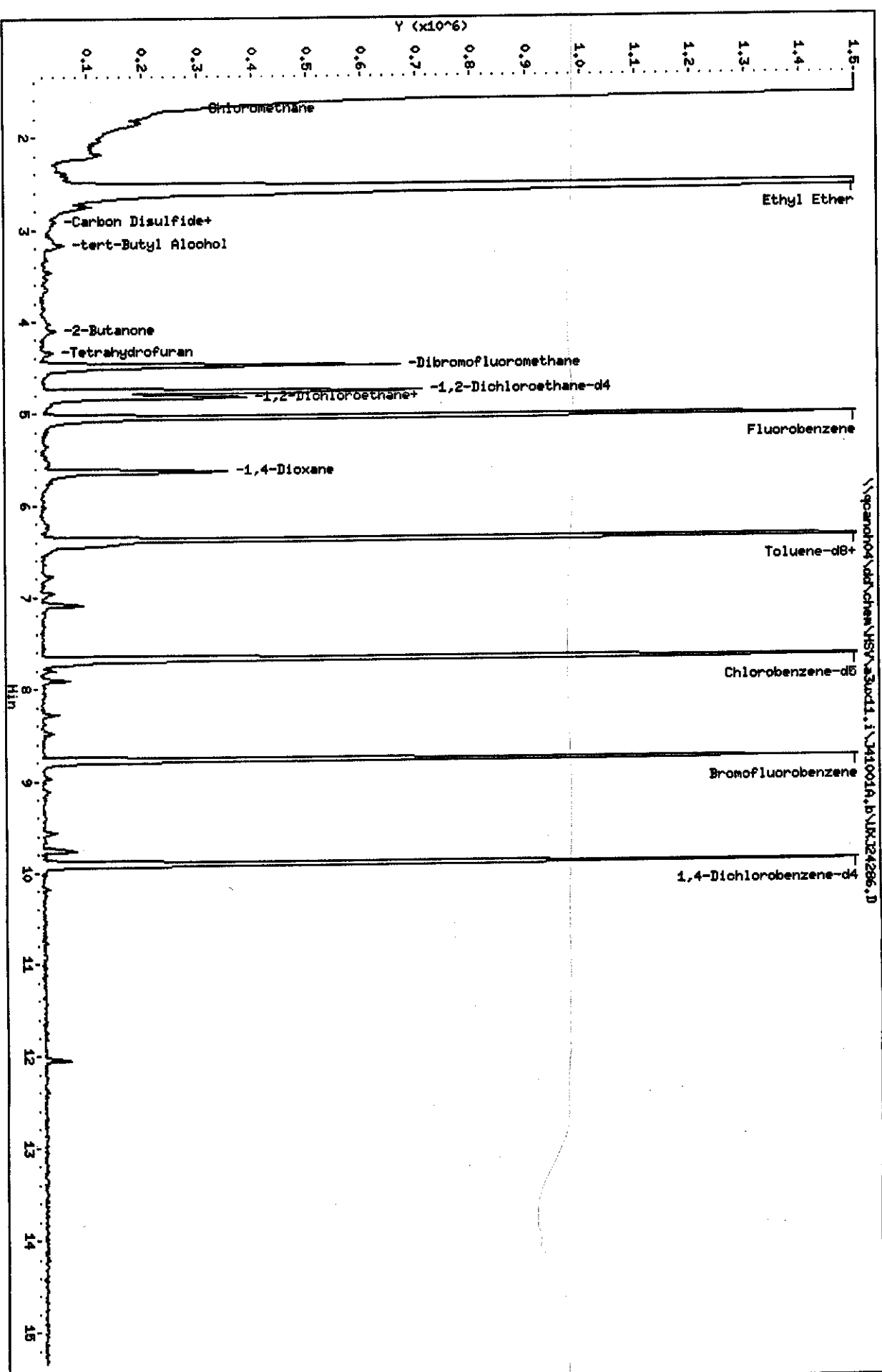
Column Phase: JPC24

Instrument: a30x11.i

Operator: 43582

Column diameter: 0.18

\\pcapn04\\dd\\chem\\NSV\\a30x11.i\\J41001A.b\\JXJ24286.D



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24286.D
Report Date: 04-Oct-2004 10:00

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24286.D
Lab Smp Id: GRDVE1AA Client Smp ID: VE540/21-26/092804
Inj Date : 01-OCT-2004 11:26
Operator : 43582 Inst ID: a3ux11.i
Smp Info : GRDVE1AA, 5ML/5ML
Misc Info : J41001A, 8260LLUX11, , 43582
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 10
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 4-8260+IX.sub
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|----------------------------|-----------|----------------|------------------------|--------|---------|----------|-----------|---------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) | (ug/L) |
| * 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | | 1775894 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | | 1562768 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | | 690306 | 50.0000 | | |
| \$ 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | | 488707 | 59.1057 | 11.821 | |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | | 634669 | 55.8214 | 11.164 | |
| \$ 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | | 1746237 | 46.5119 | 9.302 | |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | | 666906 | 41.8068 | 8.361 | |
| 8 Dichlorodifluoromethane | 85 | | Compound Not Detected. | | | | | |
| 9 Chloromethane | 50 | 1.692 | 1.704 (0.336) | | 35573 | 2.14594 | 0.4292 | |
| 10 Vinyl Chloride | 62 | | Compound Not Detected. | | | | | |
| 11 Bromomethane | 94 | | Compound Not Detected. | | | | | |
| 12 Chloroethane | 64 | | Compound Not Detected. | | | | | |
| 13 Trichlorofluoromethane | 101 | | Compound Not Detected. | | | | | |
| 15 Acrolein | 56 | | Compound Not Detected. | | | | | |
| 16 Acetone | 43 | 2.745 | 2.745 (0.545) | | 124634 | 27.4606 | 5.492 (H) | |
| 17 1,1-Dichloroethene | 96 | | Compound Not Detected. | | | | | |
| 18 Freon-113 | 151 | | Compound Not Detected. | | | | | |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | |
|---------------------------------|-----------|----------------|-------|---------------|--------|------------------------|------------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) FINAL (ug/L) |
| 19 Iodomethane | 142 | | | | | Compound Not Detected. | |
| 20 Carbon Disulfide | 76 | | 2.911 | 2.923 (0.578) | | 30113 | 1.01506 0.2030 |
| 21 Methylene Chloride | 84 | | | | | Compound Not Detected. | |
| 22 Acetonitrile | 41 | | | | | Compound Not Detected. | |
| 23 Acrylonitrile | 53 | | | | | Compound Not Detected. | |
| 24 Methyl tert-butyl ether | 73 | | | | | Compound Not Detected. | |
| 25 trans-1,2-Dichloroethene | 96 | | | | | Compound Not Detected. | |
| 26 Hexane | 86 | | | | | Compound Not Detected. | |
| 27 Vinyl acetate | 43 | | | | | Compound Not Detected. | |
| 28 1,1-Dichloroethane | 63 | | | | | Compound Not Detected. | |
| 29 tert-Butyl Alcohol | 59 | | 3.171 | 3.171 (0.629) | | 36960 | 53.5075 10.702 |
| 30 2-Butanone | 43 | | 4.106 | 4.106 (0.815) | | 56325 | 11.5796 2.316 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | | Compound Not Detected. | |
| 32 cis-1,2-dichloroethene | 96 | | | | | Compound Not Detected. | |
| 33 2,2-Dichloropropane | 77 | | | | | Compound Not Detected. | |
| 34 Bromochloromethane | 128 | | | | | Compound Not Detected. | |
| 35 Chloroform | 83 | | | | | Compound Not Detected. | |
| 36 Tetrahydrofuran | 42 | | 4.343 | 4.343 (0.862) | | 11247 | 4.29875 0.8598 |
| 37 1,1,1-Trichloroethane | 97 | | | | | Compound Not Detected. | |
| 38 1,1-Dichloropropene | 75 | | | | | Compound Not Detected. | |
| 39 Carbon Tetrachloride | 117 | | | | | Compound Not Detected. | |
| 40 1,2-Dichloroethane | 62 | | 4.828 | 4.816 (0.958) | | 314460 | 22.2762 4.455 |
| 41 Benzene | 78 | | 4.828 | 4.828 (0.958) | | 55954 | 1.36852 0.2737 |
| 42 Trichloroethene | 130 | | | | | Compound Not Detected. | |
| 43 1,2-Dichloropropane | 63 | | | | | Compound Not Detected. | |
| 44 1,4-Dioxane | 88 | | 5.633 | 5.633 (1.117) | | 309922 | 3584.83 716.96 (A) |
| 45 Dibromomethane | 93 | | | | | Compound Not Detected. | |
| 46 Bromodichloromethane | 83 | | | | | Compound Not Detected. | |
| 47 2-Chloroethyl vinyl ether | 63 | | | | | Compound Not Detected. | |
| 48 cis-1,3-Dichloropropene | 75 | | | | | Compound Not Detected. | |
| 49 4-Methyl-2-pentanone | 43 | | | | | Compound Not Detected. | |
| 50 Toluene | 91 | | 6.437 | 6.437 (0.838) | | 77426 | 1.65891 0.3318 |
| 51 trans-1,3-Dichloropropene | 75 | | | | | Compound Not Detected. | |
| 52 Ethyl Methacrylate | 69 | | | | | Compound Not Detected. | |
| 53 1,1,2-Trichloroethane | 97 | | | | | Compound Not Detected. | |
| 54 1,3-Dichloropropane | 76 | | | | | Compound Not Detected. | |
| 55 Tetrachloroethene | 164 | | | | | Compound Not Detected. | |
| 56 2-Hexanone | 43 | | | | | Compound Not Detected. | |
| 57 Dibromochloromethane | 129 | | | | | Compound Not Detected. | |
| 58 1,2-Dibromoethane | 107 | | | | | Compound Not Detected. | |
| 59 Chlorobenzene | 112 | | | | | Compound Not Detected. | |
| 60 1,1,1,2-Tetrachloroethane | 131 | | | | | Compound Not Detected. | |
| 61 Ethylbenzene | 106 | | | | | Compound Not Detected. | |
| 62 m + p-Xylene | 106 | | | | | Compound Not Detected. | |
| M 63 Xlenes (total) | 106 | | | | | Compound Not Detected. | |
| 64 Xylene-o | 106 | | | | | Compound Not Detected. | |
| 65 Styrene | 104 | | | | | Compound Not Detected. | |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|--------------------------------|-----------|------|----------------|---------------|---------|------------------------|------------|
| | | | RT | EXP RT | REL RT | RESPONSE | |
| 66 Bromoform | | 173 | | | | Compound Not Detected. | |
| 67 Isopropylbenzene | | 105 | | | | Compound Not Detected. | |
| 68 1,1,2,2-Tetrachloroethane | | 83 | | | | Compound Not Detected. | |
| 69 1,4-Dichloro-2-butene | | 53 | | | | Compound Not Detected. | |
| 70 1,2,3-Trichloropropane | | 110 | | | | Compound Not Detected. | |
| 71 Bromobenzene | | 156 | | | | Compound Not Detected. | |
| 72 n-Propylbenzene | | 120 | | | | Compound Not Detected. | |
| 73 2-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 74 1,3,5-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 75 4-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 76 tert-Butylbenzene | | 119 | | | | Compound Not Detected. | |
| 77 1,2,4-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 78 sec-Butylbenzene | | 105 | | | | Compound Not Detected. | |
| 79 4-Isopropyltoluene | | 119 | | | | Compound Not Detected. | |
| 80 1,3-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 81 1,4-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 82 n-Butylbenzene | | 91 | | | | Compound Not Detected. | |
| 83 1,2-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 84 1,2-Dibromo-3-chloropropane | | 157 | | | | Compound Not Detected. | |
| 85 1,2,4-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 86 Hexachlorobutadiene | | 225 | | | | Compound Not Detected. | |
| 87 Naphthalene | | 128 | | | | Compound Not Detected. | |
| 88 1,2,3-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 14 Dichlorofluoromethane | | 67 | | | | Compound Not Detected. | |
| 89 Ethyl Ether | | 59 | 2.544 | 2.532 (0.505) | 7937948 | 906.501 | 181.30 (A) |
| 91 3-Chloropropene | | 76 | | | | Compound Not Detected. | |
| 92 Isopropyl Ether | | 87 | | | | Compound Not Detected. | |
| 93 2-Chloro-1,3-butadiene | | 53 | | | | Compound Not Detected. | |
| 94 Propionitrile | | 54 | | | | Compound Not Detected. | |
| 95 Ethyl Acetate | | 43 | | | | Compound Not Detected. | |
| 96 Methacrylonitrile | | 41 | | | | Compound Not Detected. | |
| 97 Isobutanol | | 41 | | | | Compound Not Detected. | |
| 99 n-Butanol | | 56 | | | | Compound Not Detected. | |
| 100 Methyl Methacrylate | | 41 | | | | Compound Not Detected. | |
| 101 2-Nitropropane | | 41 | | | | Compound Not Detected. | |
| 103 Cyclohexanone | | 55 | | | | Compound Not Detected. | |
| 98 Cyclohexane | | 56 | | | | Compound Not Detected. | |
| 143 Methyl Acetate | | 43 | | | | Compound Not Detected. | |
| 144 Methylcyclohexane | | 83 | | | | Compound Not Detected. | |
| 141 1,3,5-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |

QC Flag Legend

- A - Target compound detected but, quantitated amount exceeded maximum amount.
- H - Operator selected an alternate compound hit.

Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: s3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

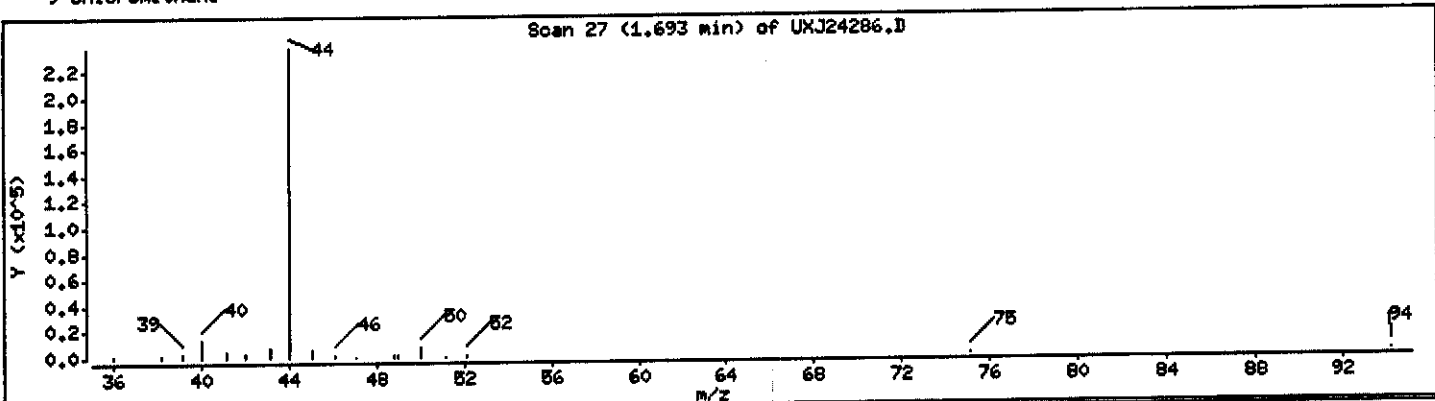
Column diameter: 0.18

Column phase: DB624

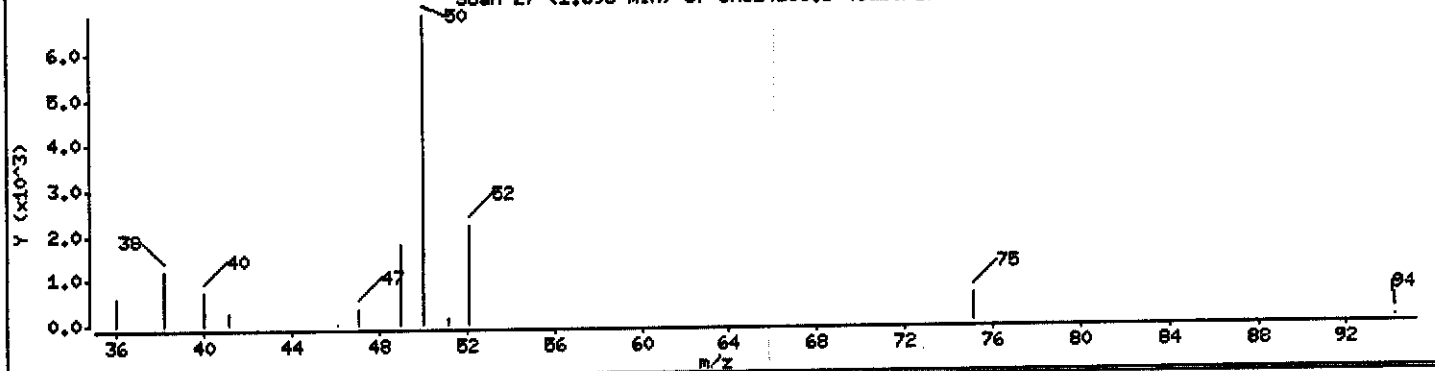
Concentration: 0.4292 ug/L

9 Chloromethane

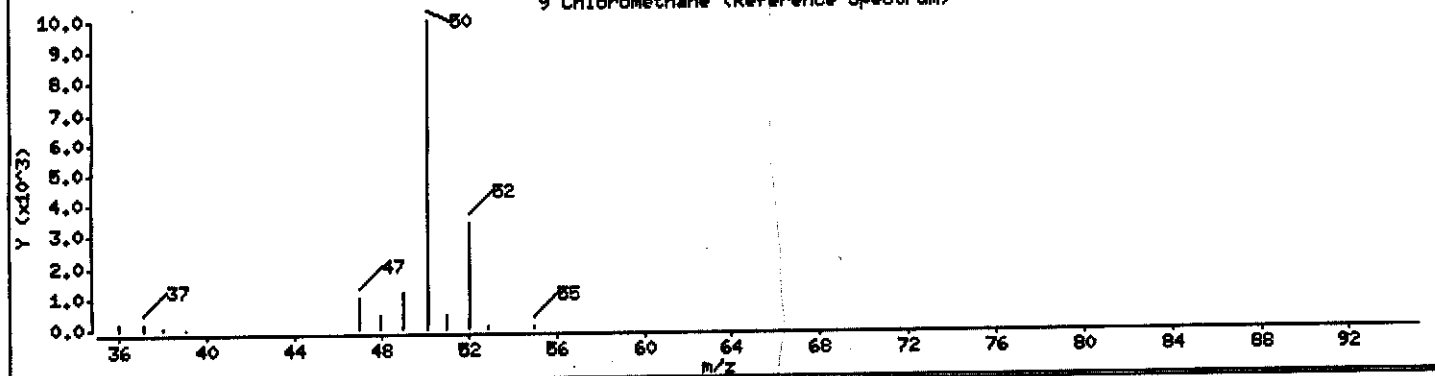
Scan 27 (1.693 min) of UXJ24286.D



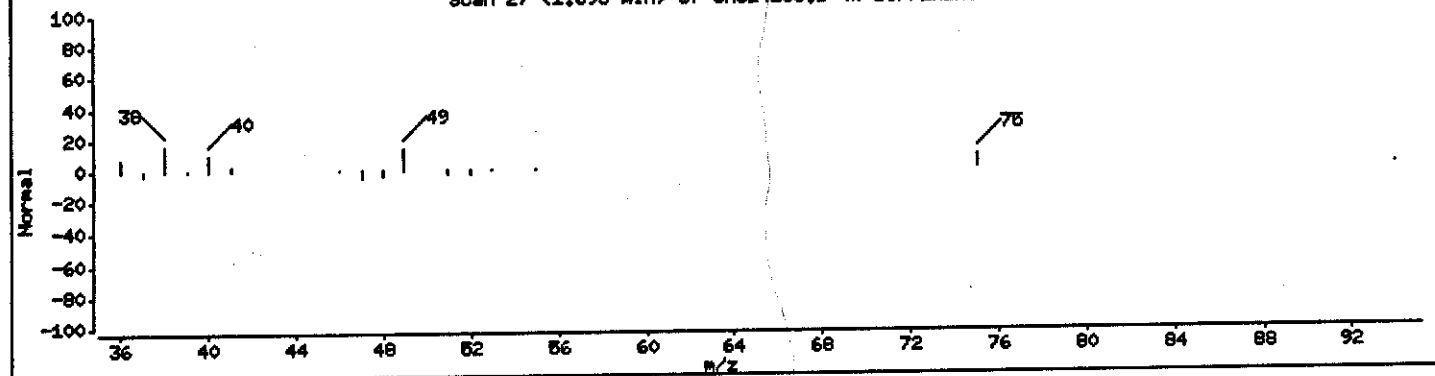
Scan 27 (1.693 min) of UXJ24286.D (Subtracted)



9 Chloromethane (Reference Spectrum)



Scan 27 (1.693 min) of UXJ24286.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: WEB40/21-26/092804

Instrument: z3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

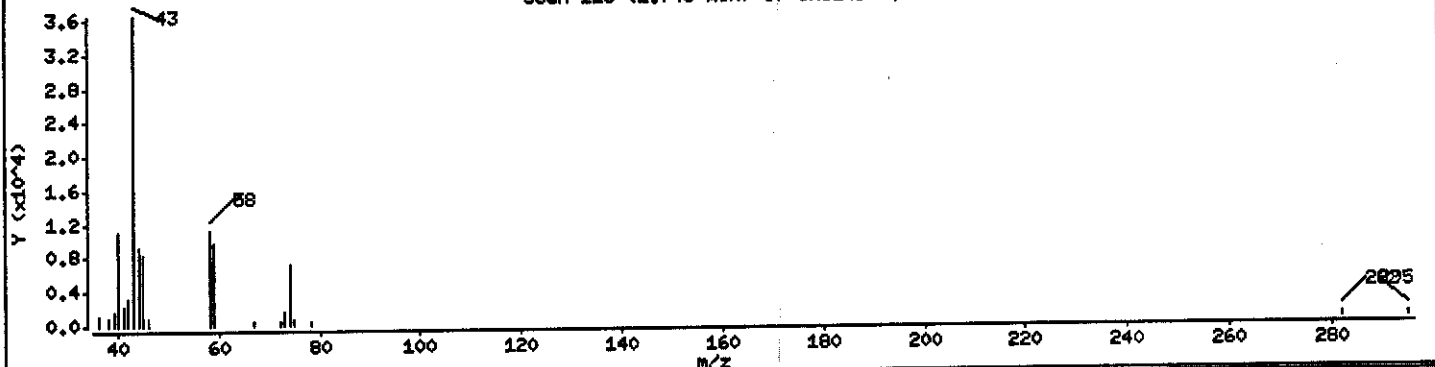
Column diameter: 0.18

Column phase: DB624

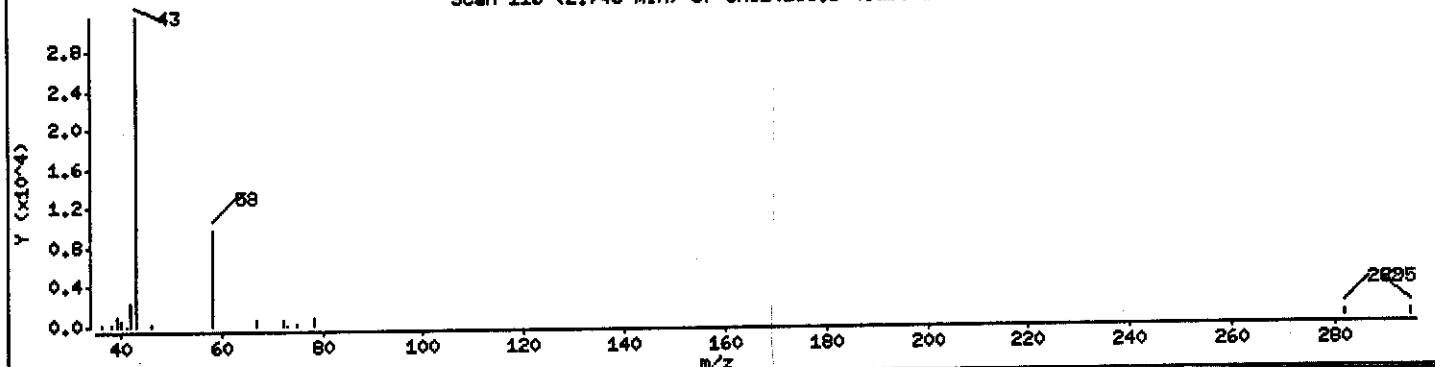
Concentration: 5.492 ug/L

16 Acetone

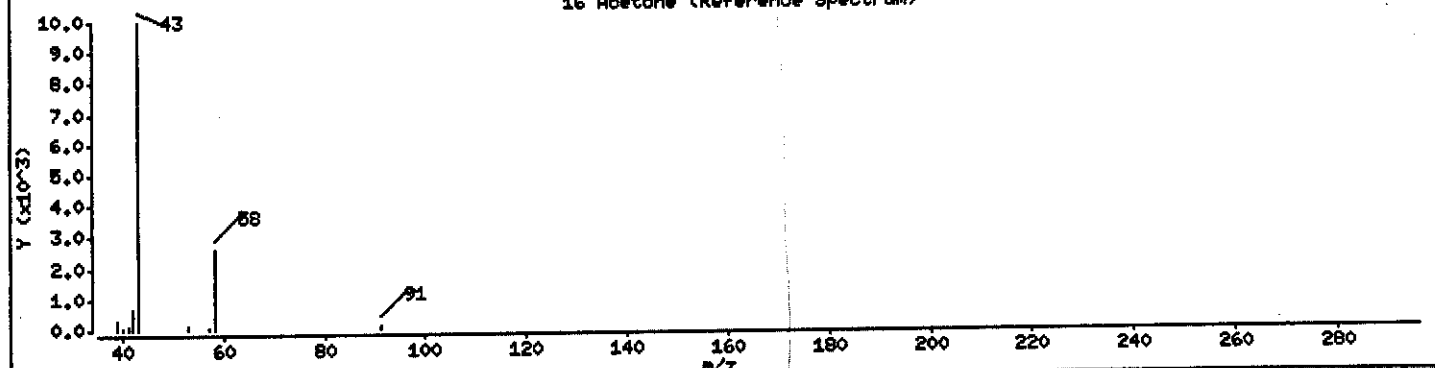
Scan 116 (2.746 min) of UXJ24286.D



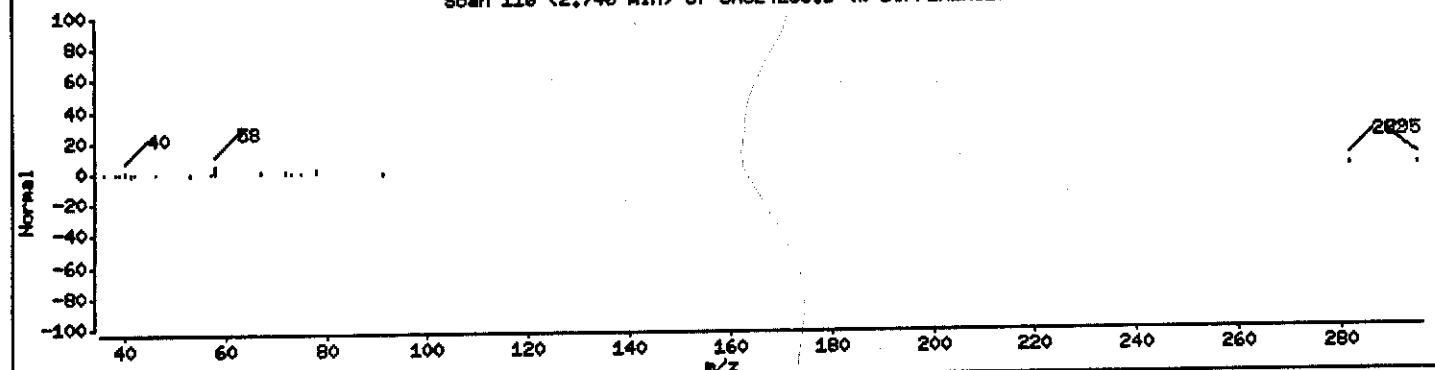
Scan 116 (2.746 min) of UXJ24286.D (Subtracted)



16 Acetone (Reference Spectrum)



Scan 116 (2.746 min) of UXJ24286.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: WEB40/21-26/092804

Instrument: z3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Purge Volume: 5.0

Operator: 43592

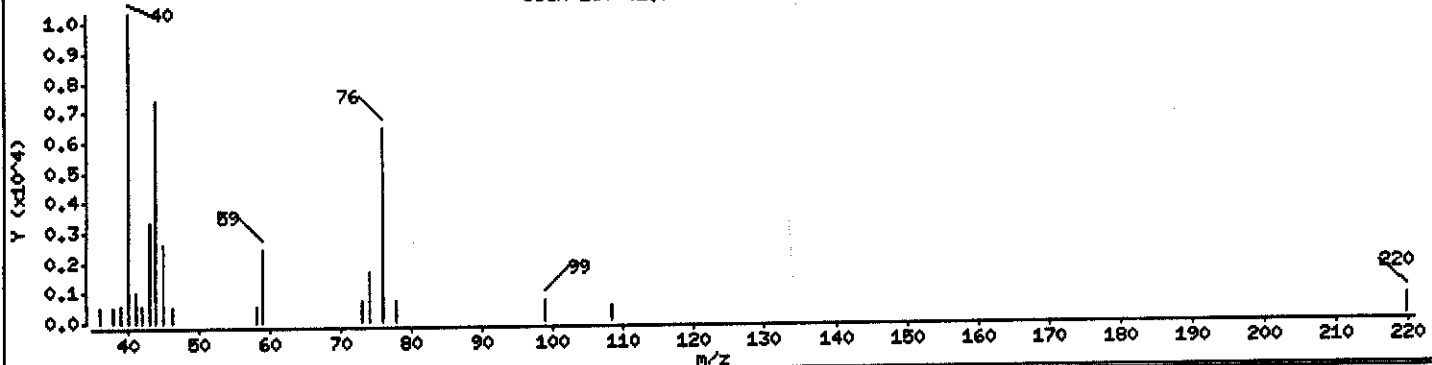
Column phase: DB624

Column diameter: 0.18

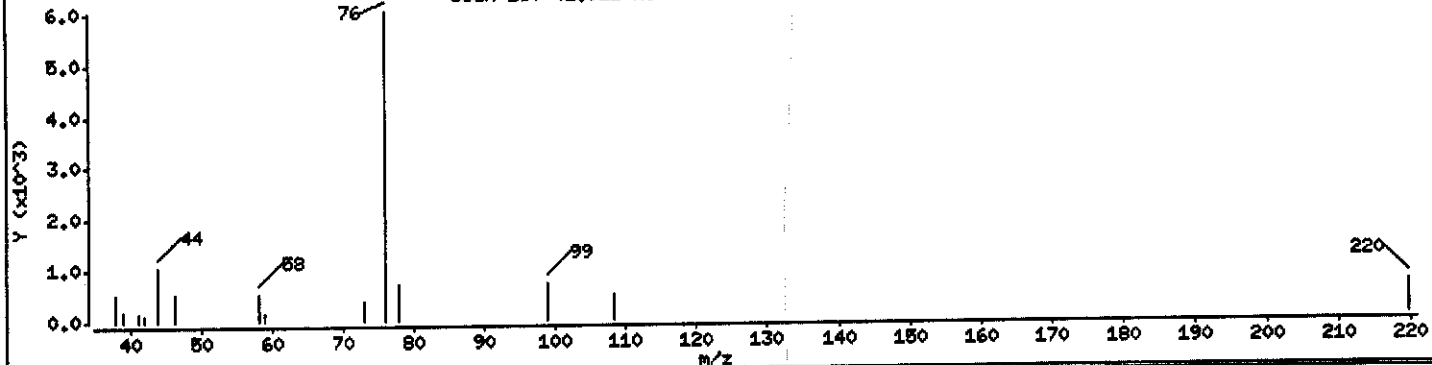
20 Carbon Disulfide

Concentration: 0.2030 ug/L

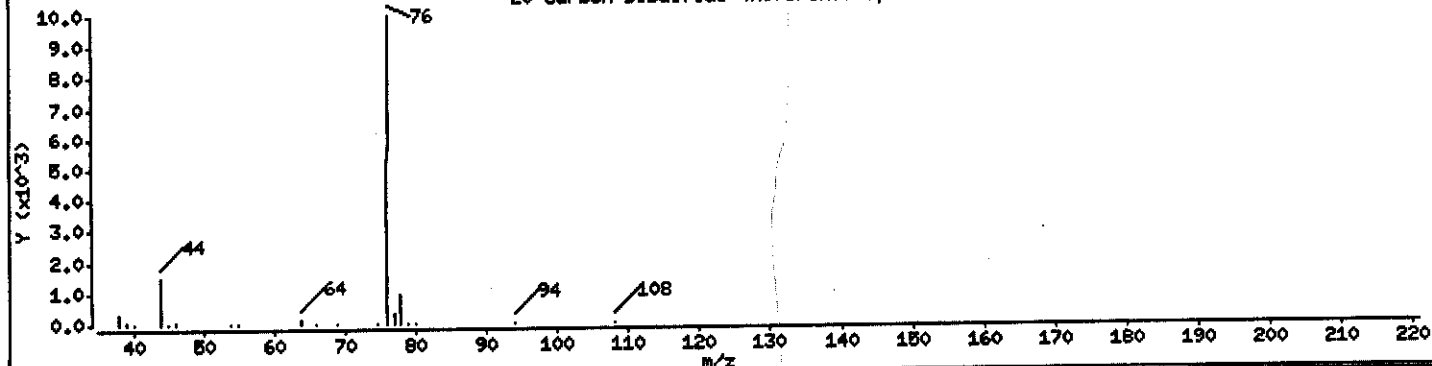
Scan 130 (2.912 min) of UXJ24286.D



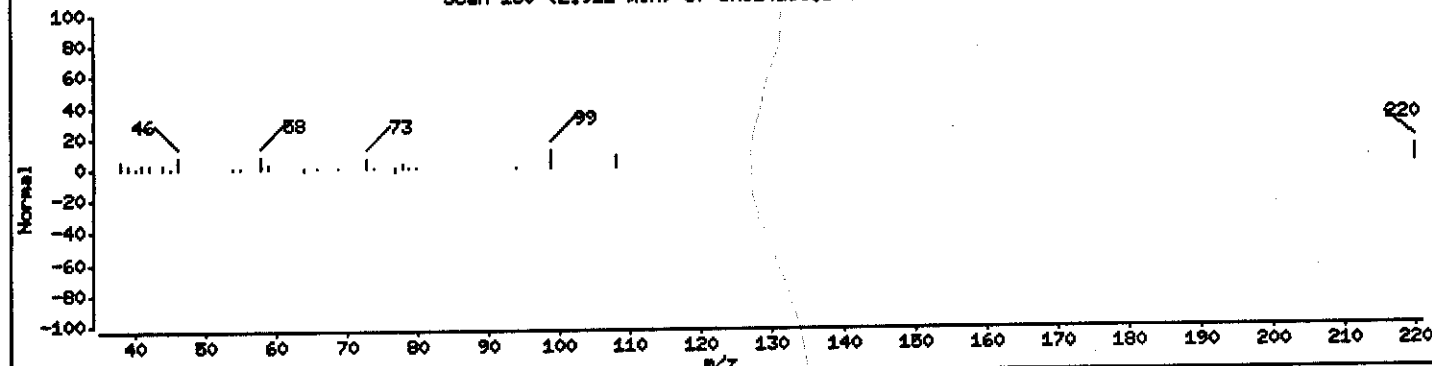
Scan 130 (2.912 min) of UXJ24286.D (Subtracted)



20 Carbon Disulfide (Reference Spectrum)



Scan 130 (2.912 min) of UXJ24286.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: z3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Purge Volume: 5.0

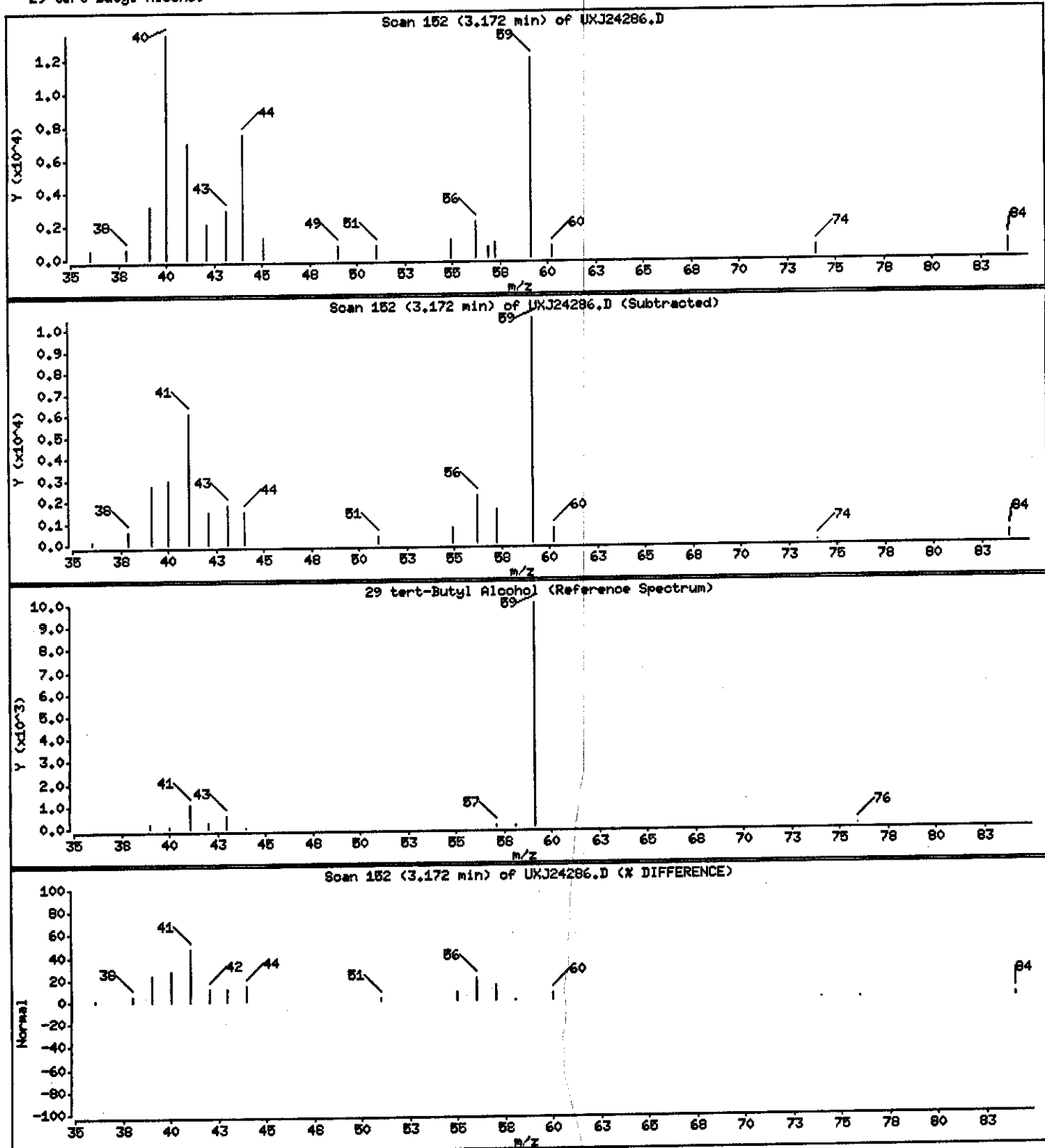
Operator: 43582

Column phaset DB624

Column diameter: 0.16

29 tert-Butyl Alcohol

Concentration: 10.702 ug/L



Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: s3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Operator: 43562

Purge Volume: 5.0

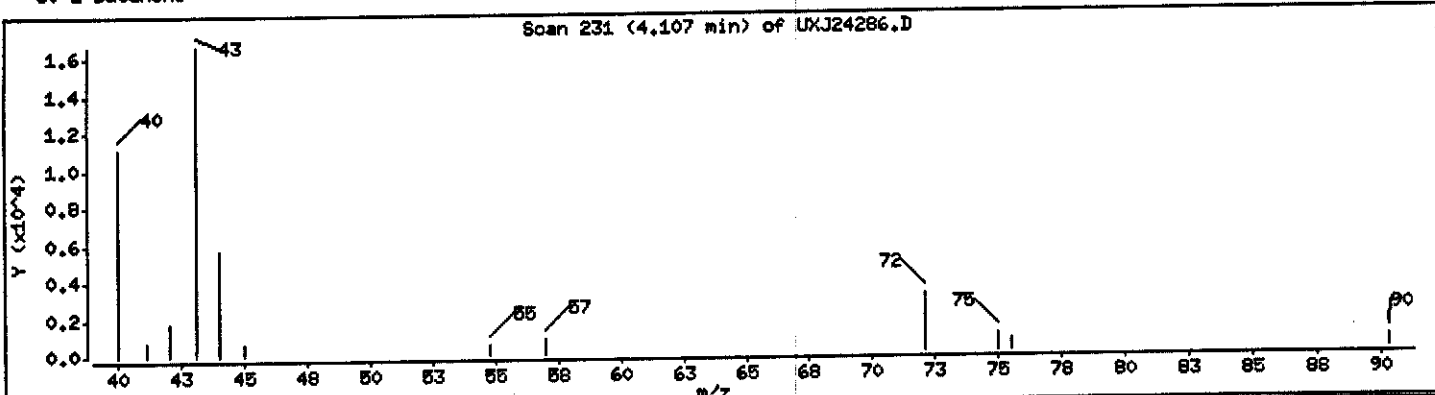
Column diameter: 0.18

Column phase: DB624

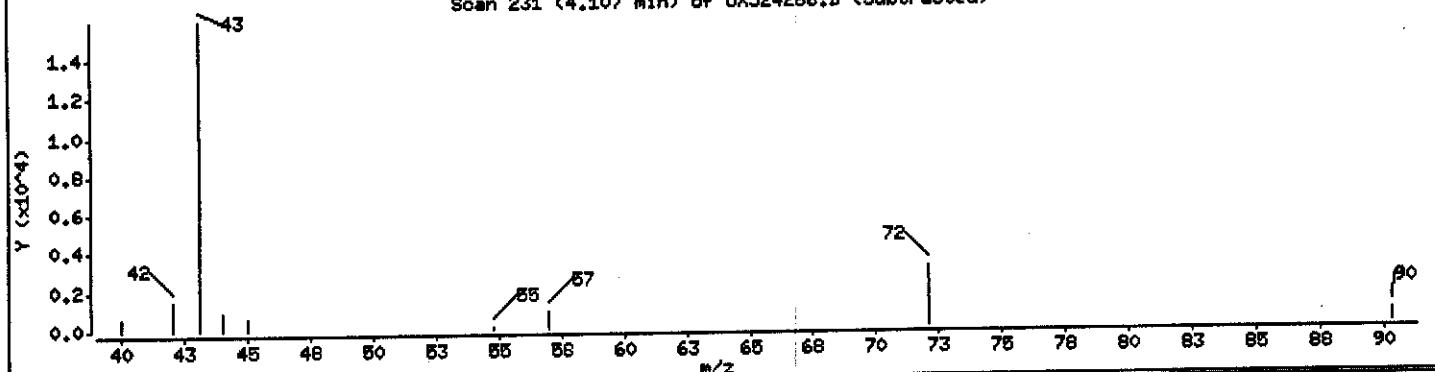
Concentration: 2.316 ug/L

30 2-Butanone

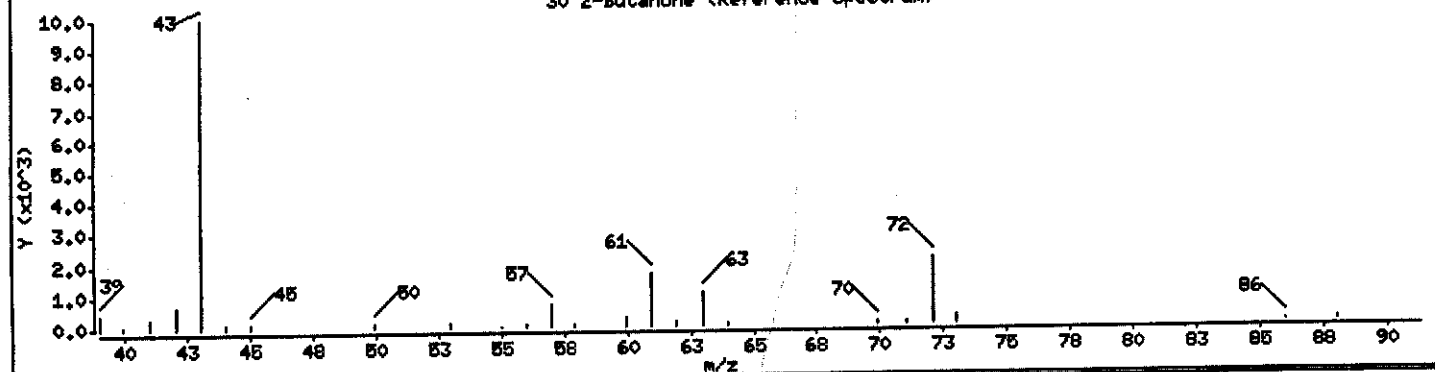
Scan 231 (4.107 min) of UXJ24286.D



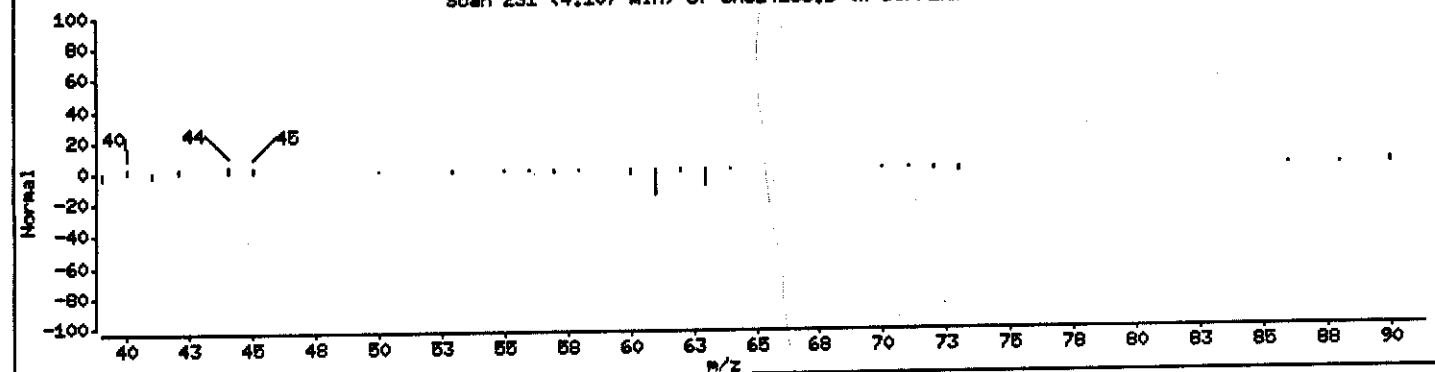
Scan 231 (4.107 min) of UXJ24286.D (Subtracted)



30 2-Butanone (Reference Spectrum)



Scan 231 (4.107 min) of UXJ24286.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: s3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

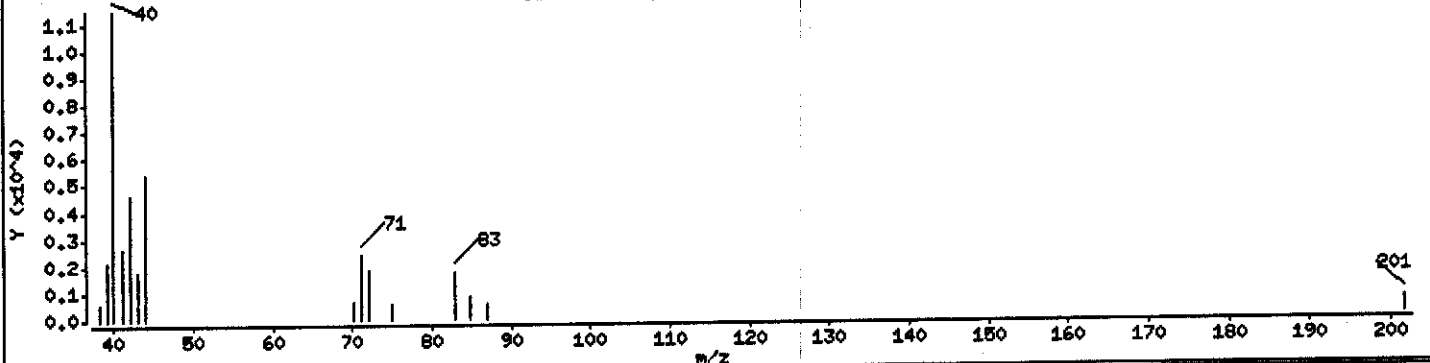
Column diameter: 0.18

Column phase: DB624

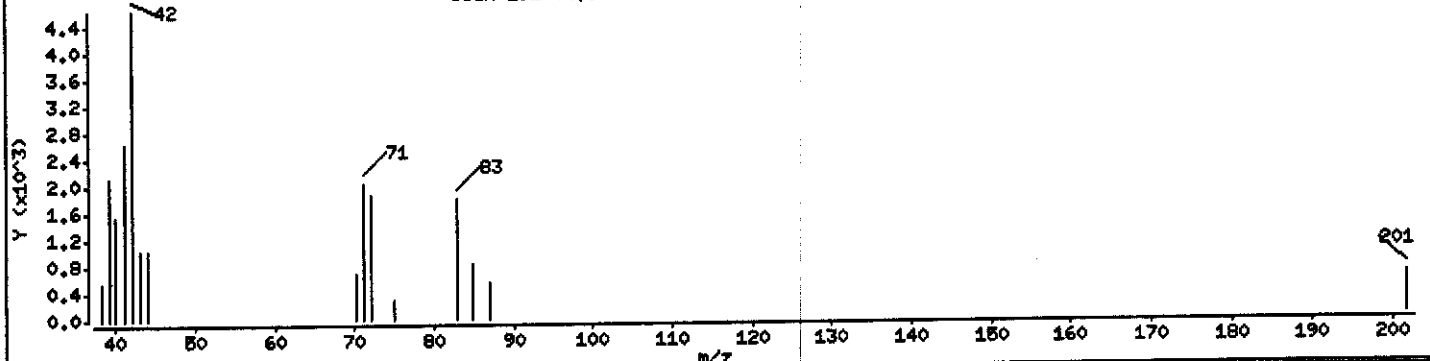
Concentration: 0.8598 ug/L

36 Tetrahydrofuran

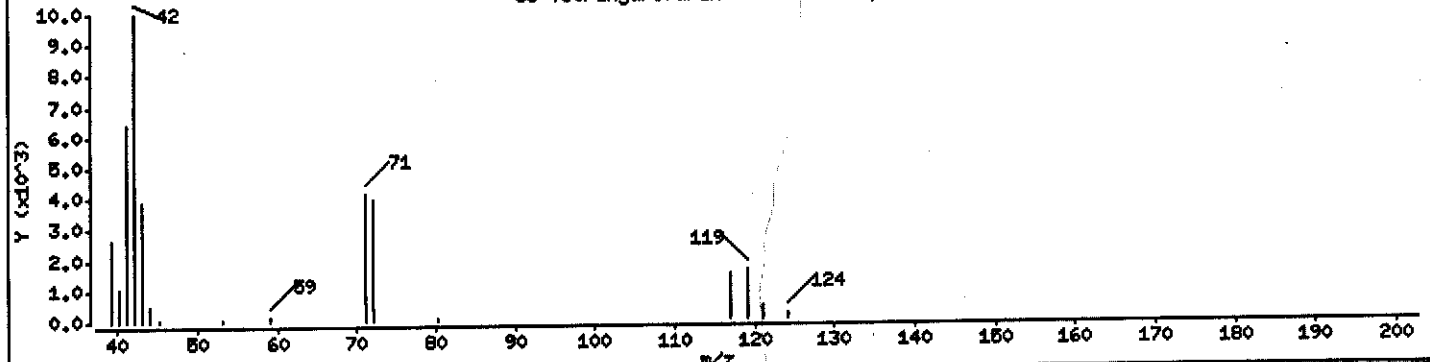
Scan 281 (4.343 min) of UXJ24286.D



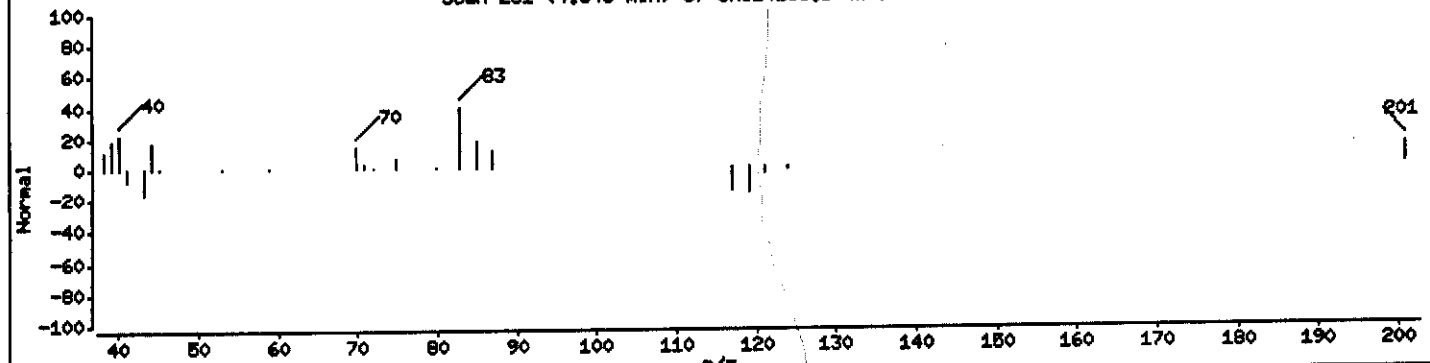
Scan 281 (4.343 min) of UXJ24286.D (Subtracted)



36 Tetrahydrofuran (Reference Spectrum)



Scan 281 (4.343 min) of UXJ24286.D (% DIFFERENCE)



Data File: \\qcanch04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: s3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

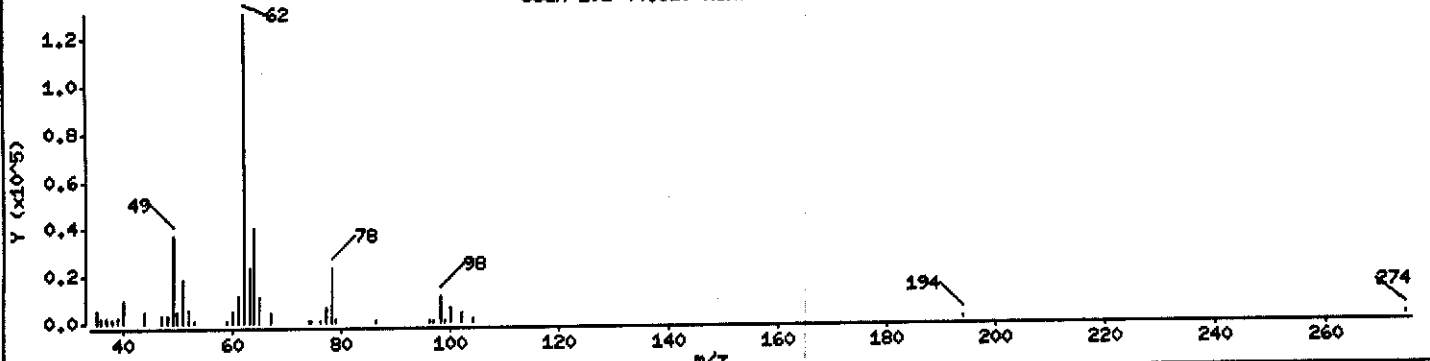
Column phase: DB624

Column diameter: 0.18

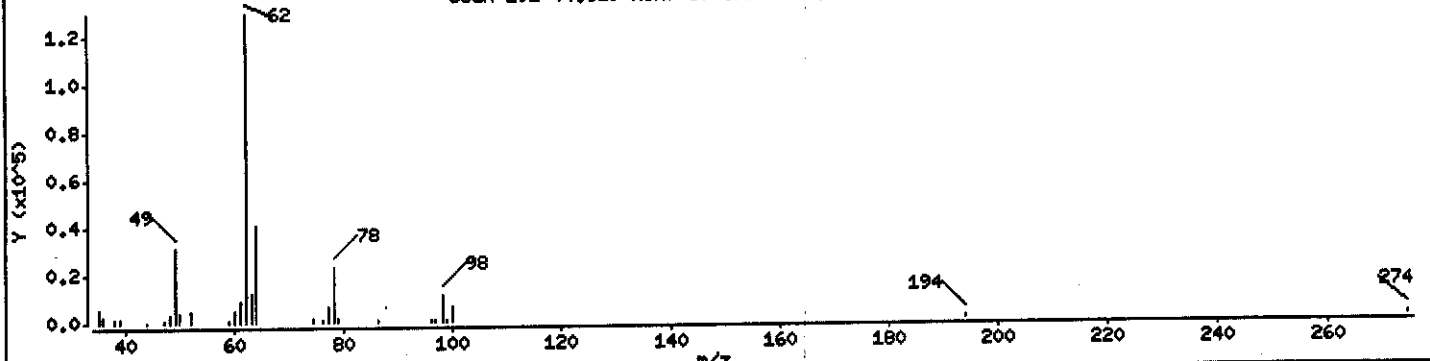
40 1,2-Dichloroethane

Concentration: 4.465 ug/L

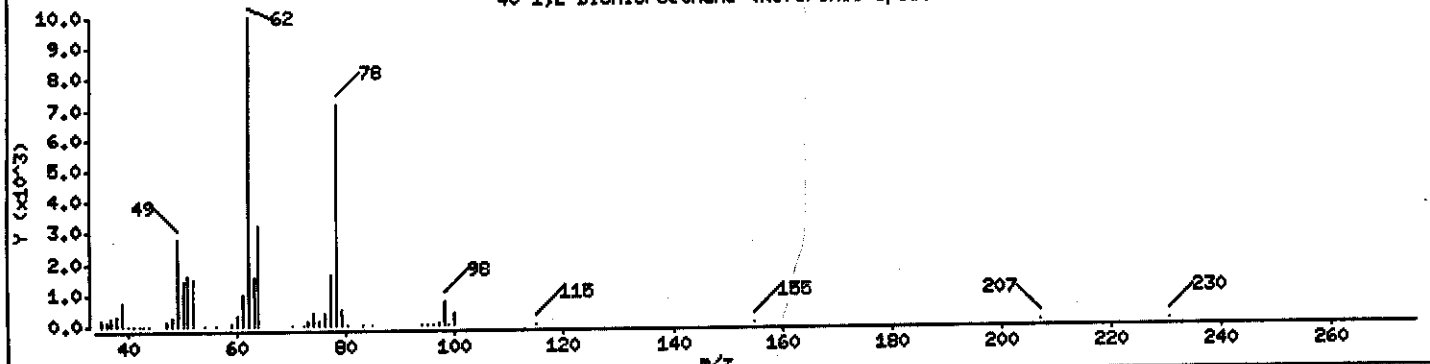
Scan 292 (4.829 min) of UXJ24286.D



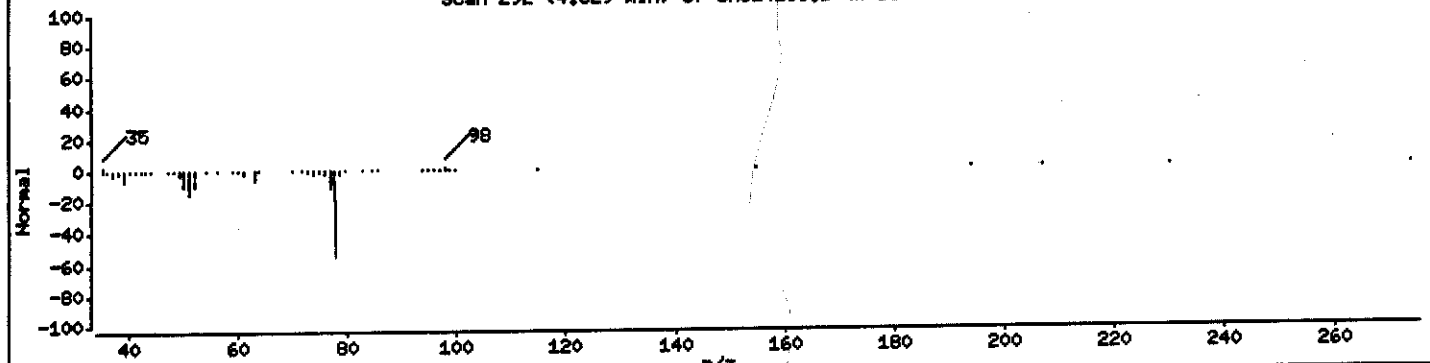
Scan 292 (4.829 min) of UXJ24286.D (Subtracted)



40 1,2-Dichloroethane (Reference Spectrum)



Scan 292 (4.829 min) of UXJ24286.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VEB40/21-26/092804

Instrument: z3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Purge Volume: 5.0

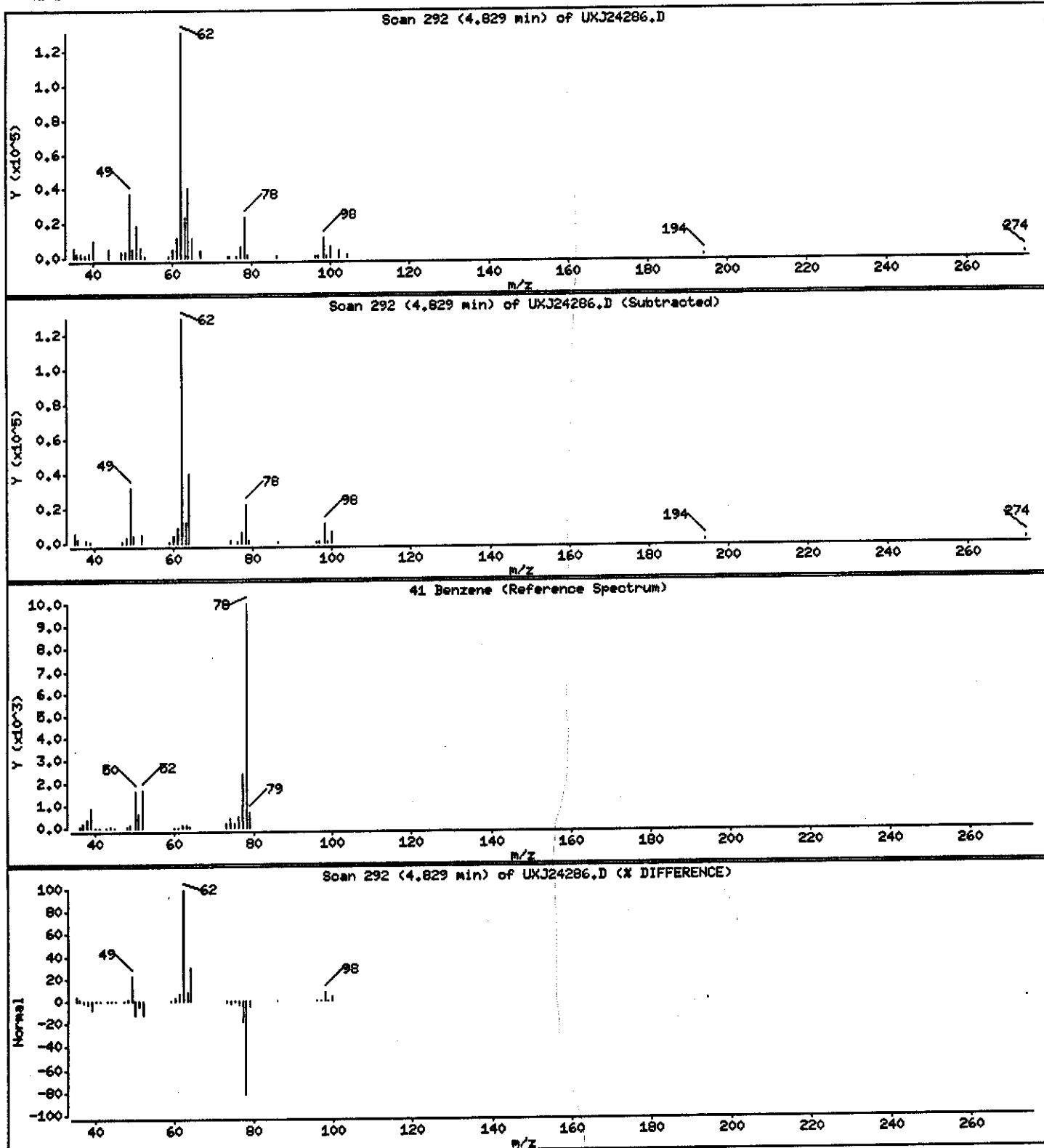
Operator: 43582

Column phase: DB624

Column diameter: 0.18

41 Benzene

Concentration: 0.2737 ug/L



Data File: \\qcanoh04\dd\chem\HSV\z3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: z3ux11.i

Sample Info: DRIVE1AA,5ML/5ML

Operator: 43582

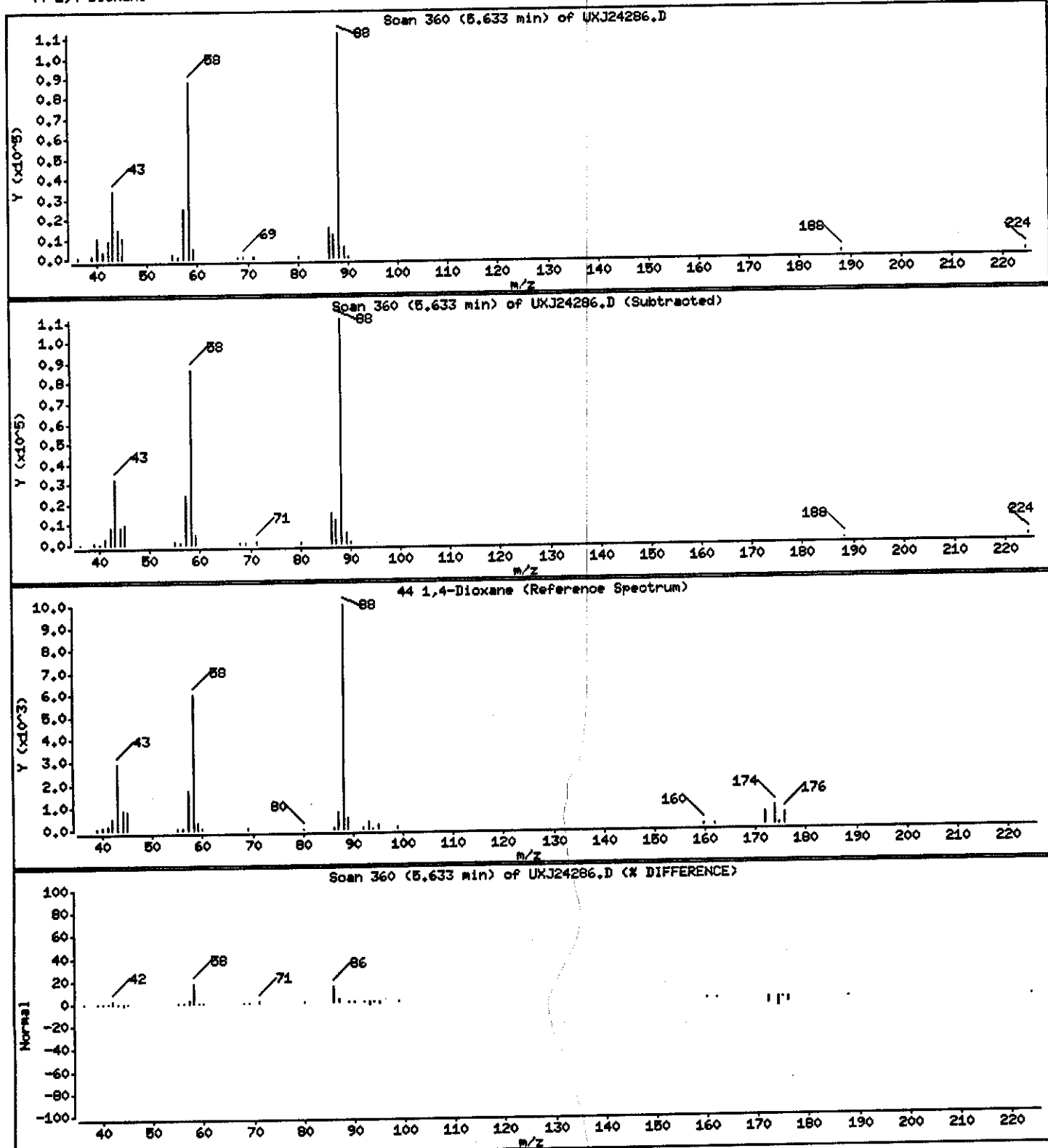
Purge Volume: 5.0

Column diameter: 0.18

Column phase: DB624

Concentration: 716.96 ug/L

44 1,4-Dioxane



Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: s3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Purge Volume: 5.0

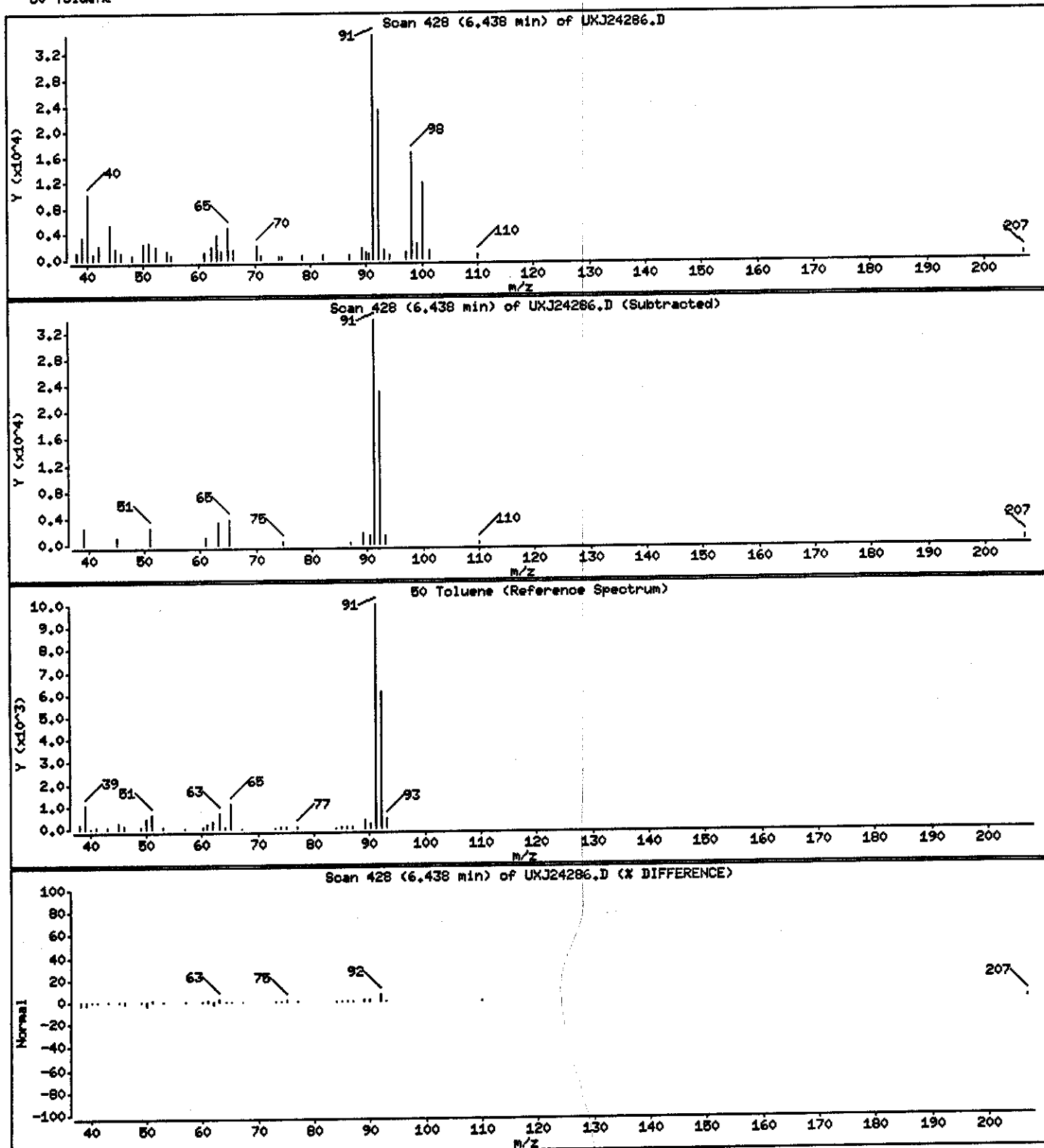
Operator: 43582

Column phase: DB624

Column diameter: 0.18

50 Toluene

Concentration: 0.3318 ug/L



Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24286.D

Date : 01-OCT-2004 11:26

Client ID: VE540/21-26/092804

Instrument: s3ux11.i

Sample Info: GRDVE1AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

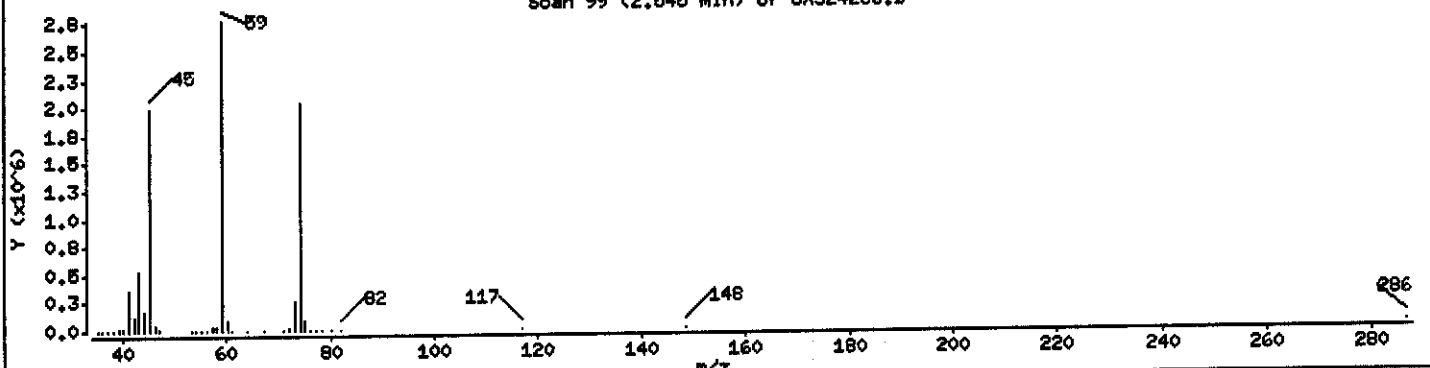
Column diameter: 0.18

Column phase: DB624

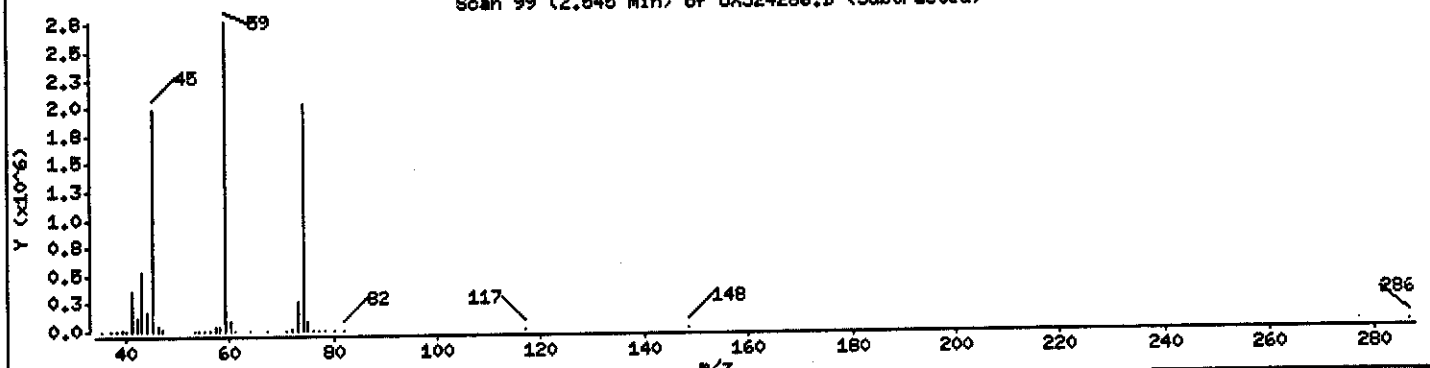
Concentration: 181.30 ug/L

89 Ethyl Ether

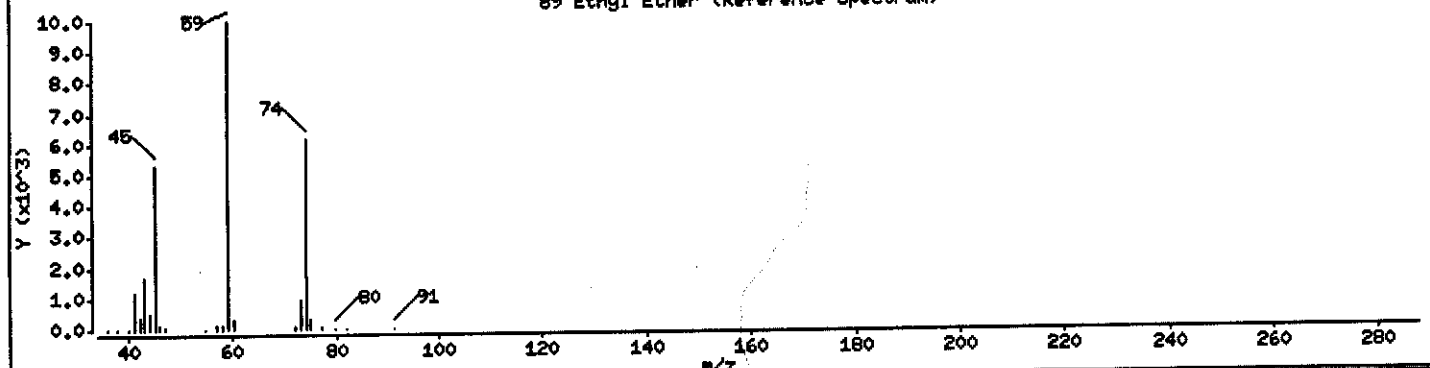
Scan 99 (2.545 min) of UXJ24286.D



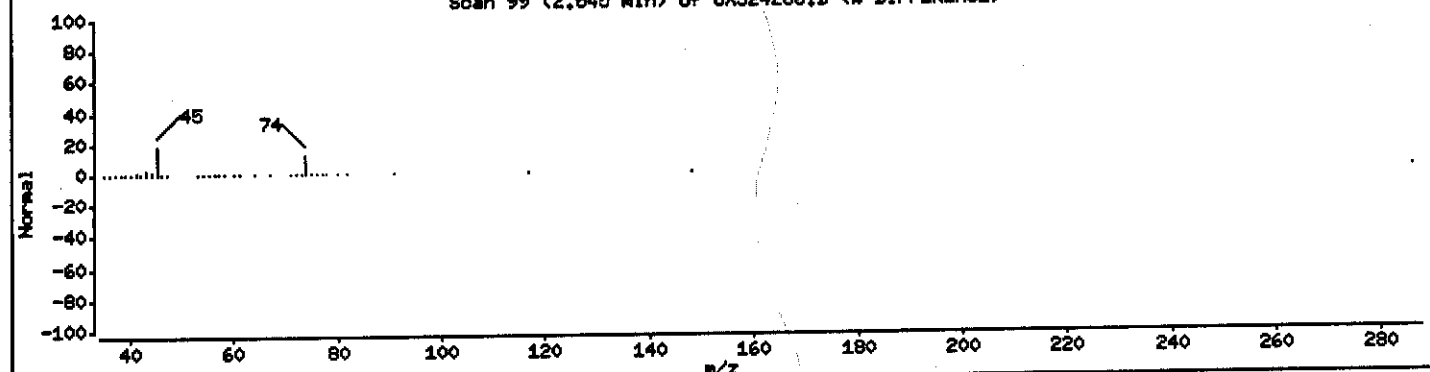
Scan 99 (2.545 min) of UXJ24286.D (Subtracted)



89 Ethyl Ether (Reference Spectrum)



Scan 99 (2.545 min) of UXJ24286.D (% DIFFERENCE)



PAYNE FIRM INC.

Client Sample ID: VE540/35.5-40.5/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-002 Work Order #....: GRDX11AA Matrix.....: WG
 Date Sampled....: 09/28/04 13:30 Date Received...: 09/29/04
 Prep Date.....: 10/01/04 Analysis Date...: 10/01/04
 Prep Batch #....: 4275213
 Dilution Factor: 1 Initial Wgt/Vol: 5 mL Final Wgt/Vol.: 5 mL
 Method.....: SW846 8260B

| PARAMETER | RESULT | REPORTING | |
|------------------------------------|----------|-----------|-------|
| | | LIMIT | UNITS |
| Acetone | 6.8 J | 10 | ug/L |
| Acetonitrile | ND | 20 | ug/L |
| Acrolein | ND | 20 | ug/L |
| Acrylonitrile | ND | 20 | ug/L |
| Benzene | ND | 1.0 | ug/L |
| Bromodichloromethane | ND | 1.0 | ug/L |
| Bromoform | ND | 1.0 | ug/L |
| Bromomethane | ND | 1.0 | ug/L |
| 2-Butanone | 1.7 J | 10 | ug/L |
| Carbon disulfide | ND | 1.0 | ug/L |
| Carbon tetrachloride | ND | 1.0 | ug/L |
| Chlorobenzene | ND | 1.0 | ug/L |
| Chloroprene | ND | 2.0 | ug/L |
| Dibromochloromethane | ND | 1.0 | ug/L |
| Chloroethane | ND | 1.0 | ug/L |
| Chloroform | 0.18 J | 1.0 | ug/L |
| Chloromethane | 0.24 J,B | 1.0 | ug/L |
| 3-Chloropropene | ND | 2.0 | ug/L |
| 1,2-Dibromo-3-chloropropane (DBCP) | ND | 2.0 | ug/L |
| 1,2-Dibromoethane | ND | 1.0 | ug/L |
| Dibromomethane | ND | 1.0 | ug/L |
| trans-1,4-Dichloro-2-butene | ND | 1.0 | ug/L |
| 1,1-Dichloroethane | ND | 1.0 | ug/L |
| 1,2-Dichloroethane | ND | 1.0 | ug/L |
| cis-1,2-Dichloroethene | ND | 1.0 | ug/L |
| trans-1,2-Dichloroethene | ND | 1.0 | ug/L |
| 1,1-Dichloroethene | ND | 1.0 | ug/L |
| 1,2-Dichloroethene (total) | ND | 2.0 | ug/L |
| Dichlorofluoromethane | ND | 2.0 | ug/L |
| 1,2-Dichloropropane | ND | 1.0 | ug/L |
| cis-1,3-Dichloropropene | ND | 1.0 | ug/L |
| trans-1,3-Dichloropropene | ND | 1.0 | ug/L |
| 1,4-Dioxane | 20 J | 50 | ug/L |
| Ethylbenzene | ND | 1.0 | ug/L |
| Ethyl methacrylate | ND | 1.0 | ug/L |

(Continued on next page)

PAYNE FIRM INC.

Client Sample ID: VE540/35.5-40.5/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-002 Work Order #....: GRDX11AA Matrix.....: WG

| PARAMETER | RESULT | REPORTING | |
|---------------------------|--------|-----------|-------|
| | | LIMIT | UNITS |
| 2-Hexanone | ND | 10 | ug/L |
| Iodomethane | ND | 1.0 | ug/L |
| Isobutanol | ND | 50 | ug/L |
| Methacrylonitrile | ND | 2.0 | ug/L |
| Methylene chloride | ND | 1.0 | ug/L |
| Methyl methacrylate | ND | 2.0 | ug/L |
| 4-Methyl-2-pentanone | ND | 10 | ug/L |
| Propionitrile | ND | 4.0 | ug/L |
| Styrene | ND | 1.0 | ug/L |
| 1,1,1,2-Tetrachloroethane | ND | 1.0 | ug/L |
| 1,1,2,2-Tetrachloroethane | ND | 1.0 | ug/L |
| Tetrachloroethene | ND | 1.0 | ug/L |
| Toluene | ND | 1.0 | ug/L |
| 1,1,1-Trichloroethane | 0.48 J | 1.0 | ug/L |
| 1,1,2-Trichloroethane | ND | 1.0 | ug/L |
| Trichloroethene | 0.39 J | 1.0 | ug/L |
| Trichlorofluoromethane | ND | 1.0 | ug/L |
| 1,2,3-Trichloropropane | ND | 1.0 | ug/L |
| Vinyl acetate | ND | 2.0 | ug/L |
| Vinyl chloride | ND | 1.0 | ug/L |
| Xylenes (total) | ND | 2.0 | ug/L |

| SURROGATE | PERCENT RECOVERY | RECOVERY | |
|-----------------------|---------------------|----------|--------|
| | | LIMITS | |
| Dibromofluoromethane | 119 | (73 | - 122) |
| 1,2-Dichloroethane-d4 | 113 | (61 | - 128) |
| Toluene-d8 | 90 | (76 | - 110) |
| 4-Bromofluorobenzene | 81 | (74 | - 116) |

NOTE(S) :

J Estimated result. Result is less than RL.

B Method blank contamination. The associated method blank contains the target analyte at a reportable level.

Data File: \\spando04\\data\\chem\\HSI\\a3dx11.i\\J41001A.b\\JMK24287.D

Date : 01-OCT-2004 11:49

Client ID: 4E546735.5-40.5.092

Sample Info: GRDX10A.BHL-FML

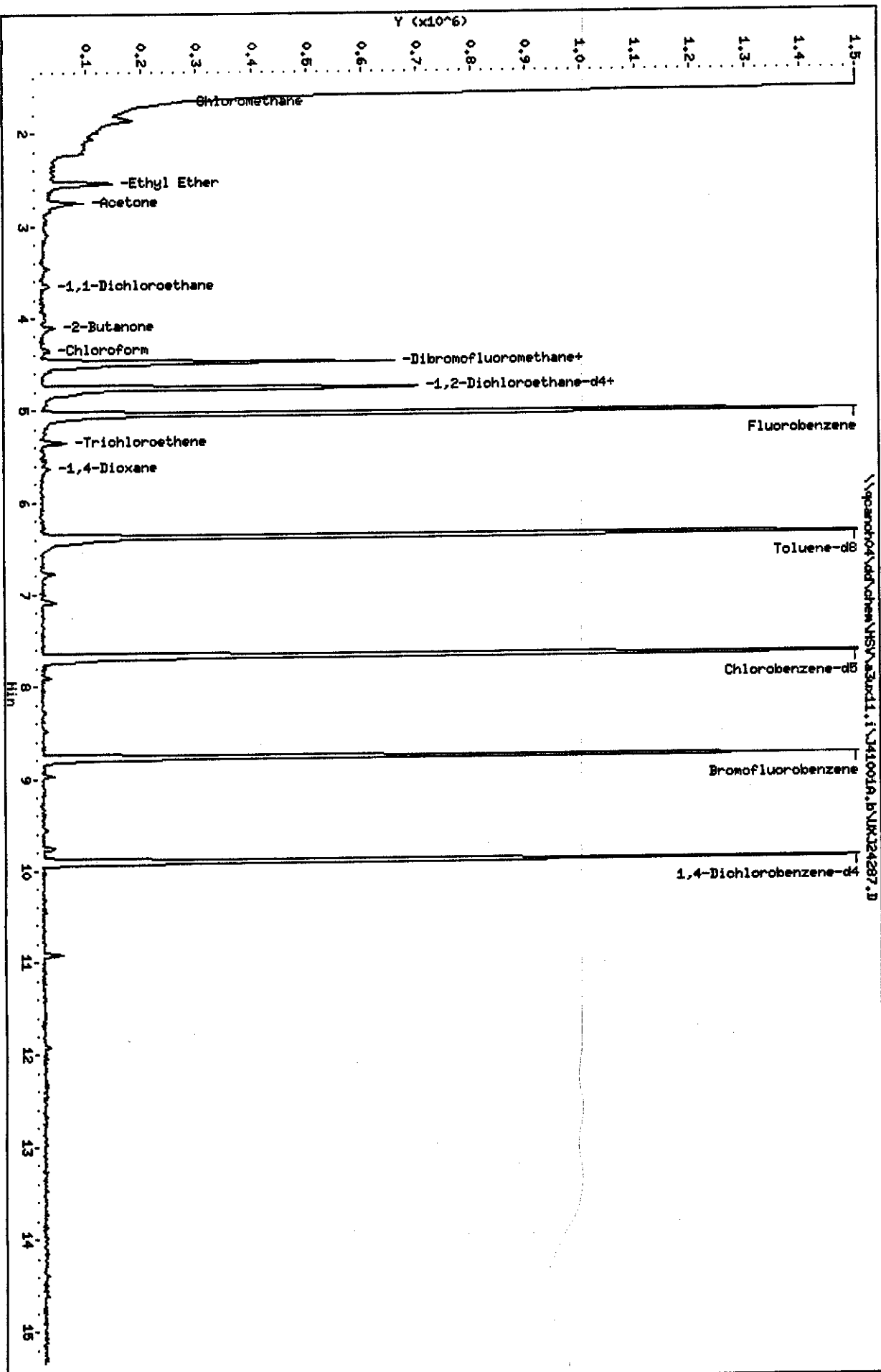
Purge Volume: 5.0

Column Phase: Bk24

Instrument: a3dx11.i

Operator: 43682

Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24287.D
Report Date: 04-Oct-2004 10:00

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24287.D
Lab Smp Id: GRDX11AA Client Smp ID: VE540/35.5-40.5/092
Inj Date : 01-OCT-2004 11:49
Operator : 43582 Inst ID: a3ux11.i
Smp Info : GRDX11AA, 5ML/5ML
Misc Info : J41001A, 8260LLUX11, , 43582
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 11
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 4-8260+IX.sub
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|-----------|--------------------------|----------------|-------|------------------------|---------|----------|--------|---------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) | (ug/L) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 1755793 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1533746 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 651571 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 484626 | 59.2831 | 11.857 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 635910 | 56.5708 | 11.314 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 1660418 | 45.0629 | 9.012 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 636446 | 40.6523 | 8.130 | |
| 8 | Dichlorodifluoromethane | 85 | | Compound Not Detected. | | | | |
| 9 | Chloromethane | 50 | 1.669 | 1.704 (0.331) | 19838 | 1.21043 | 0.2421 | |
| 10 | Vinyl Chloride | 62 | | Compound Not Detected. | | | | |
| 11 | Bromomethane | 94 | | Compound Not Detected. | | | | |
| 12 | Chloroethane | 64 | | Compound Not Detected. | | | | |
| 13 | Trichlorofluoromethane | 101 | | Compound Not Detected. | | | | |
| 15 | Acrolein | 56 | | Compound Not Detected. | | | | |
| 16 | Acetone | 43 | 2.745 | 2.745 (0.545) | 148382 | 34.1914 | 6.838 | |
| 17 | 1,1-Dichloroethene | 96 | | Compound Not Detected. | | | | |
| 18 | Freon-113 | 151 | | Compound Not Detected. | | | | |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | | |
|---------------------------------|-----------|------|----------------|---------------|--------|------------------------|---------|--------|
| | | | RT | EXP RT | REL RT | RESPONSE | | |
| 19 Iodomethane | | 142 | | | | Compound Not Detected. | | |
| 20 Carbon Disulfide | | 76 | | | | Compound Not Detected. | | |
| 21 Methylene Chloride | | 84 | | | | Compound Not Detected. | | |
| 22 Acetonitrile | | 41 | | | | Compound Not Detected. | | |
| 23 Acrylonitrile | | 53 | | | | Compound Not Detected. | | |
| 24 Methyl tert-butyl ether | | 73 | | | | Compound Not Detected. | | |
| 25 trans-1,2-Dichloroethene | | 96 | | | | Compound Not Detected. | | |
| 26 Hexane | | 86 | | | | Compound Not Detected. | | |
| 27 Vinyl acetate | | 43 | | | | Compound Not Detected. | | |
| 28 1,1-Dichloroethane | | 63 | 3.657 | 3.657 (0.725) | | 14048 | 0.82673 | 0.1653 |
| 29 tert-Butyl Alcohol | | 59 | | | | Compound Not Detected. | | |
| 30 2-Butanone | | 43 | 4.106 | 4.106 (0.815) | | 39726 | 8.26058 | 1.652 |
| M 31 1,2-Dichloroethene (total) | | 96 | | | | Compound Not Detected. | | |
| 32 cis-1,2-dichloroethene | | 96 | | | | Compound Not Detected. | | |
| 33 2,2-Dichloropropane | | 77 | | | | Compound Not Detected. | | |
| 34 Bromochloromethane | | 128 | | | | Compound Not Detected. | | |
| 35 Chloroform | | 83 | 4.355 | 4.355 (0.864) | | 14984 | 0.87721 | 0.1754 |
| 36 Tetrahydrofuran | | 42 | | | | Compound Not Detected. | | |
| 37 1,1,1-Trichloroethane | | 97 | 4.520 | 4.520 (0.897) | | 30753 | 2.38385 | 0.4768 |
| 38 1,1-Dichloropropene | | 75 | | | | Compound Not Detected. | | |
| 39 Carbon Tetrachloride | | 117 | | | | Compound Not Detected. | | |
| 40 1,2-Dichloroethane | | 62 | | | | Compound Not Detected. | | |
| 41 Benzene | | 78 | 4.828 | 4.828 (0.958) | | 34577 | 0.85536 | 0.1711 |
| 42 Trichloroethene | | 130 | 5.349 | 5.349 (1.061) | | 18006 | 1.95171 | 0.3903 |
| 43 1,2-Dichloropropene | | 63 | | | | Compound Not Detected. | | |
| 44 1,4-Dioxane | | 88 | 5.644 | 5.633 (1.120) | | 8526 | 99.7482 | 19.950 |
| 45 Dibromomethane | | 93 | | | | Compound Not Detected. | | |
| 46 Bromodichloromethane | | 83 | | | | Compound Not Detected. | | |
| 47 2-Chloroethyl vinyl ether | | 63 | | | | Compound Not Detected. | | |
| 48 cis-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | | |
| 49 4-Methyl-2-pentanone | | 43 | | | | Compound Not Detected. | | |
| 50 Toluene | | 91 | | | | Compound Not Detected. | | |
| 51 trans-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | | |
| 52 Ethyl Methacrylate | | 69 | | | | Compound Not Detected. | | |
| 53 1,1,2-Trichloroethane | | 97 | | | | Compound Not Detected. | | |
| 54 1,3-Dichloropropene | | 76 | | | | Compound Not Detected. | | |
| 55 Tetrachloroethene | | 164 | | | | Compound Not Detected. | | |
| 56 2-Hexanone | | 43 | | | | Compound Not Detected. | | |
| 57 Dibromochloromethane | | 129 | | | | Compound Not Detected. | | |
| 58 1,2-Dibromoethane | | 107 | | | | Compound Not Detected. | | |
| 59 Chlorobenzene | | 112 | | | | Compound Not Detected. | | |
| 60 1,1,1,2-Tetrachloroethane | | 131 | | | | Compound Not Detected. | | |
| 61 Ethylbenzene | | 106 | | | | Compound Not Detected. | | |
| 62 m + p-Xylene | | 106 | | | | Compound Not Detected. | | |
| M 63 Xlenes (total) | | 106 | | | | Compound Not Detected. | | |
| 64 Xylene-o | | 106 | | | | Compound Not Detected. | | |
| 65 Styrene | | 104 | | | | Compound Not Detected. | | |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|--------------------------------|-----------|------|----------------|---------------|--------|------------------------|-------------------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 66 Bromoform | | 173 | | | | Compound Not Detected. | |
| 67 Isopropylbenzene | | 105 | | | | Compound Not Detected. | |
| 68 1,1,2,2-Tetrachloroethane | | 83 | | | | Compound Not Detected. | |
| 69 1,4-Dichloro-2-butene | | 53 | | | | Compound Not Detected. | |
| 70 1,2,3-Trichloropropane | | 110 | | | | Compound Not Detected. | |
| 71 Bromobenzene | | 156 | | | | Compound Not Detected. | |
| 72 n-Propylbenzene | | 120 | | | | Compound Not Detected. | |
| 73 2-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 74 1,3,5-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 75 4-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 76 tert-Butylbenzene | | 119 | | | | Compound Not Detected. | |
| 77 1,2,4-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 78 sec-Butylbenzene | | 105 | | | | Compound Not Detected. | |
| 79 4-Isopropyltoluene | | 119 | | | | Compound Not Detected. | |
| 80 1,3-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 81 1,4-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 82 n-Butylbenzene | | 91 | | | | Compound Not Detected. | |
| 83 1,2-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 84 1,2-Dibromo-3-chloropropane | | 157 | | | | Compound Not Detected. | |
| 85 1,2,4-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 86 Hexachlorobutadiene | | 225 | | | | Compound Not Detected. | |
| 87 Naphthalene | | 128 | | | | Compound Not Detected. | |
| 88 1,2,3-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 14 Dichlorofluoromethane | | 67 | | | | Compound Not Detected. | |
| 89 Ethyl Ether | | 59 | 2.544 | 2.532 (0.505) | 100480 | 11.6060 | 2.321 |
| 91 3-Chloropropene | | 76 | | | | Compound Not Detected. | |
| 92 Isopropyl Ether | | 87 | | | | Compound Not Detected. | |
| 93 2-Chloro-1,3-butadiene | | 53 | | | | Compound Not Detected. | |
| 94 Propionitrile | | 54 | | | | Compound Not Detected. | |
| 95 Ethyl Acetate | | 43 | | | | Compound Not Detected. | |
| 96 Methacrylonitrile | | 41 | | | | Compound Not Detected. | |
| 97 Isobutanol | | 41 | | | | Compound Not Detected. | |
| 99 n-Butanol | | 56 | | | | Compound Not Detected. | |
| 100 Methyl Methacrylate | | 41 | | | | Compound Not Detected. | |
| 101 2-Nitropropane | | 41 | | | | Compound Not Detected. | |
| 103 Cyclohexanone | | 55 | | | | Compound Not Detected. | |
| 98 Cyclohexane | | 56 | | | | Compound Not Detected. | |
| 143 Methyl Acetate | | 43 | | | | Compound Not Detected. | |
| 144 Methylcyclohexane | | 83 | | | | Compound Not Detected. | |
| 141 1,3,5-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |

Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

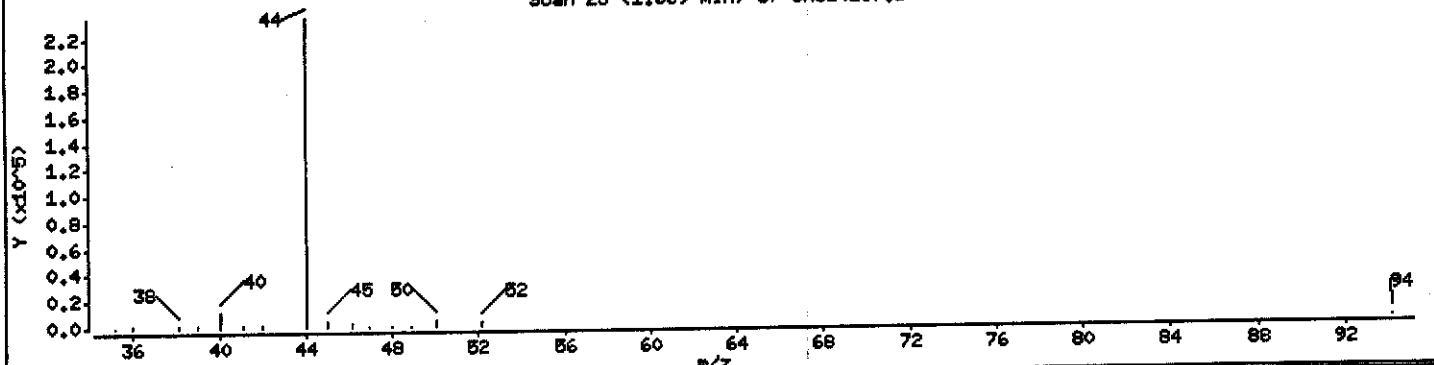
Column phase: DB624

Column diameter: 0.18

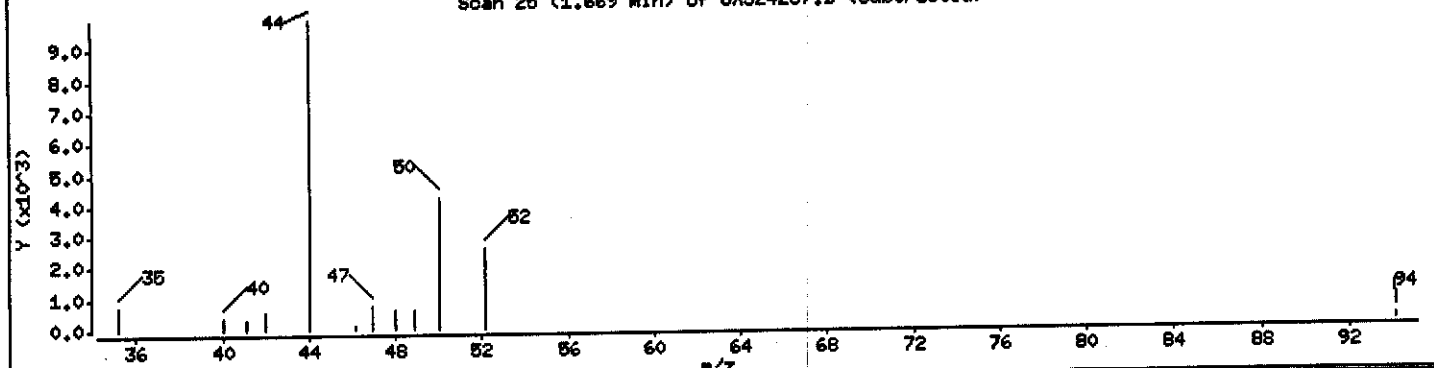
9 Chloromethane

Concentration: 0.2421 ug/L

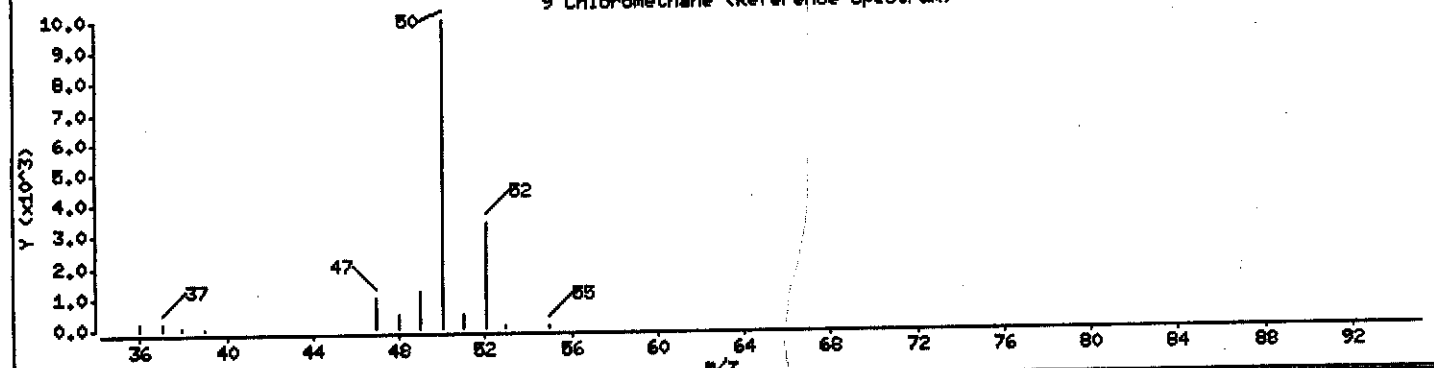
Scan 25 (1.669 min) of UXJ24287.D



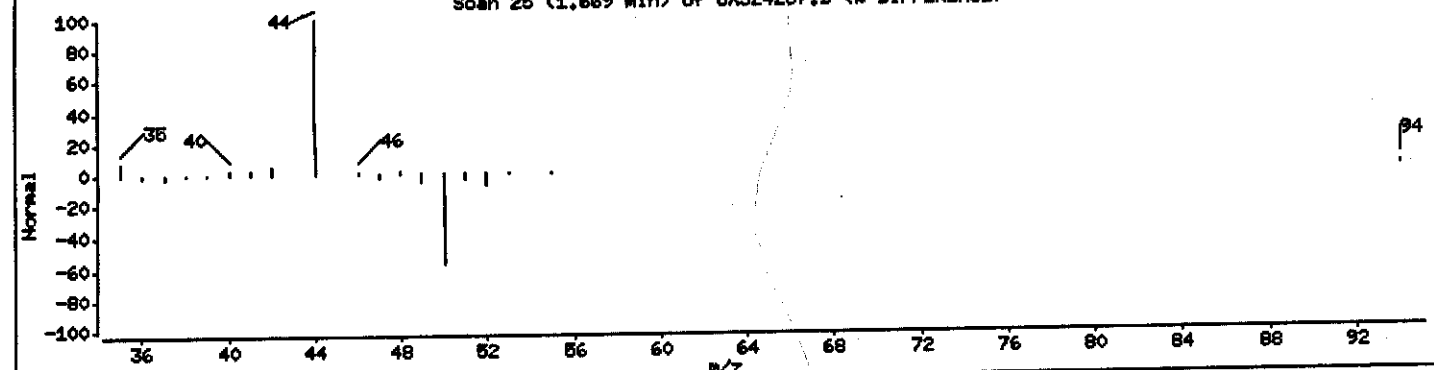
Scan 25 (1.669 min) of UXJ24287.D (Subtracted)



9 Chloromethane (Reference Spectrum)



Scan 25 (1.669 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

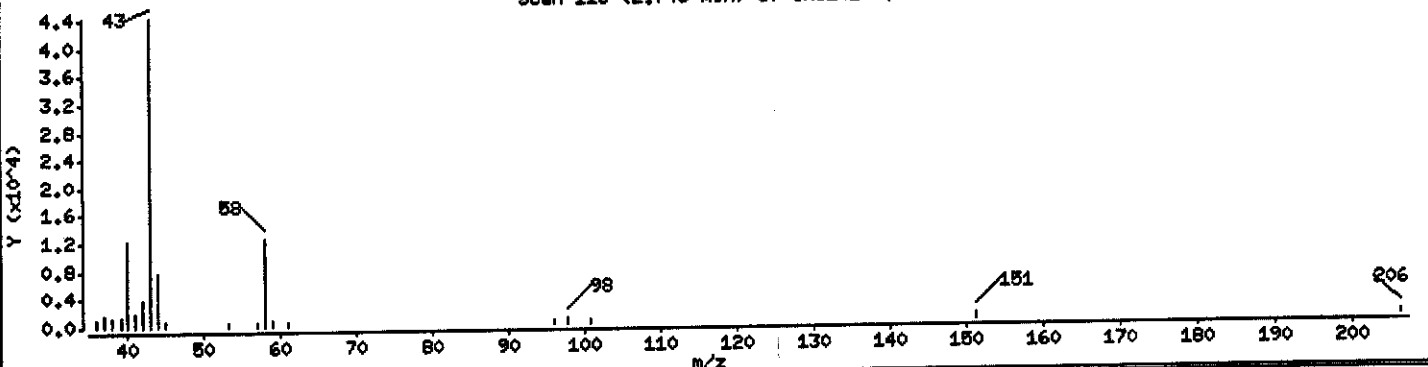
Column diameter: 0.18

Column phase: DB624

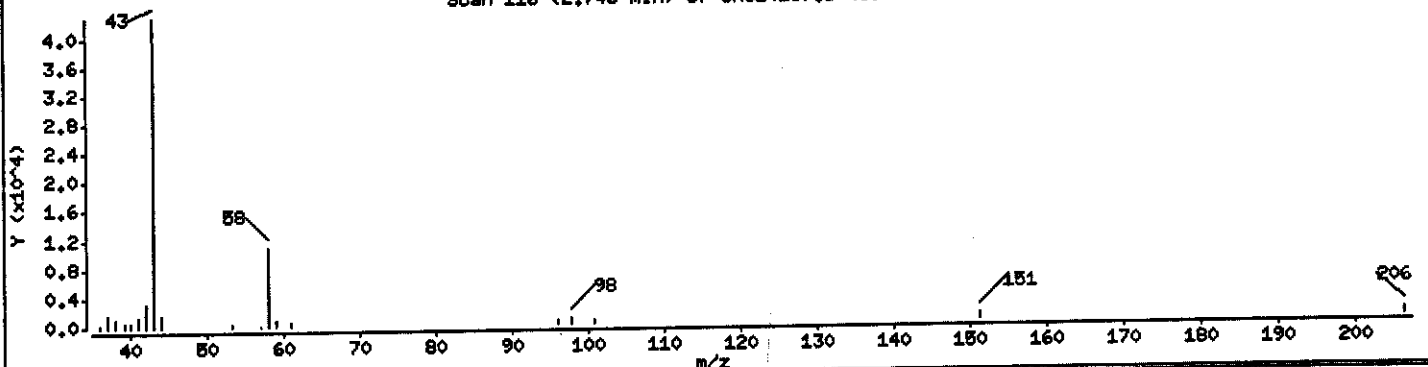
Concentration: 6.838 ug/L

16 Acetone

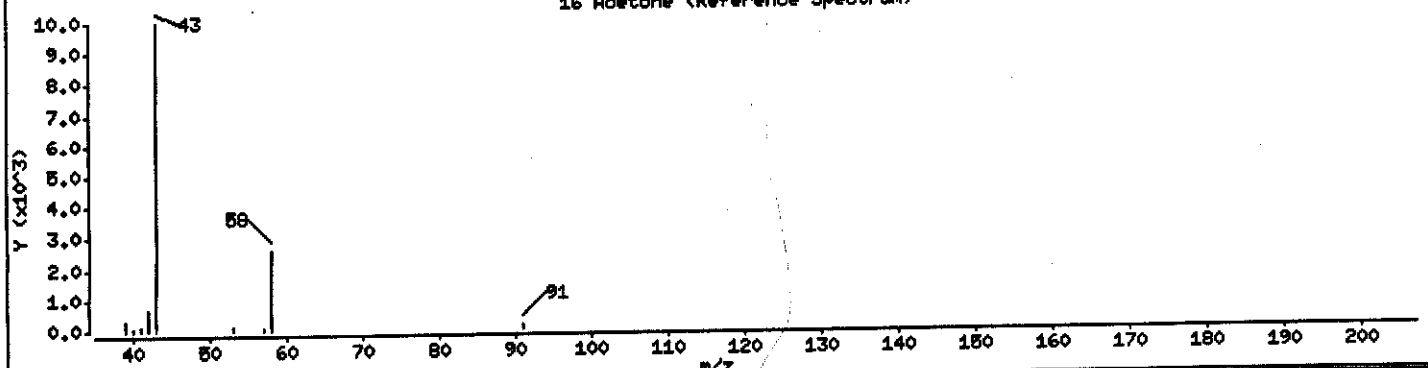
Scan 116 (2.746 min) of UXJ24287.D



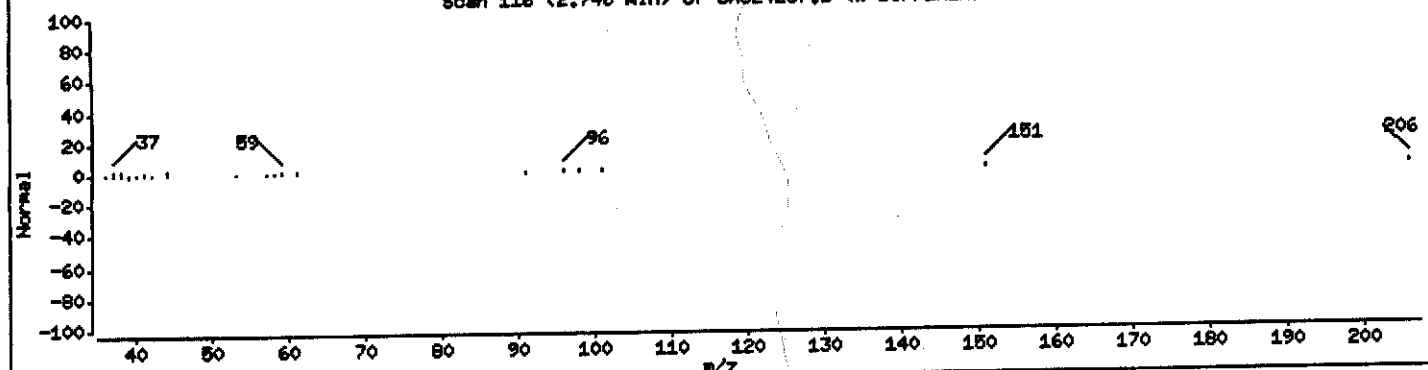
Scan 116 (2.746 min) of UXJ24287.D (Subtracted)



16 Acetone (Reference Spectrum)



Scan 116 (2.746 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qpanch04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: s3ux11.i

Sample Info: CRDX11AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

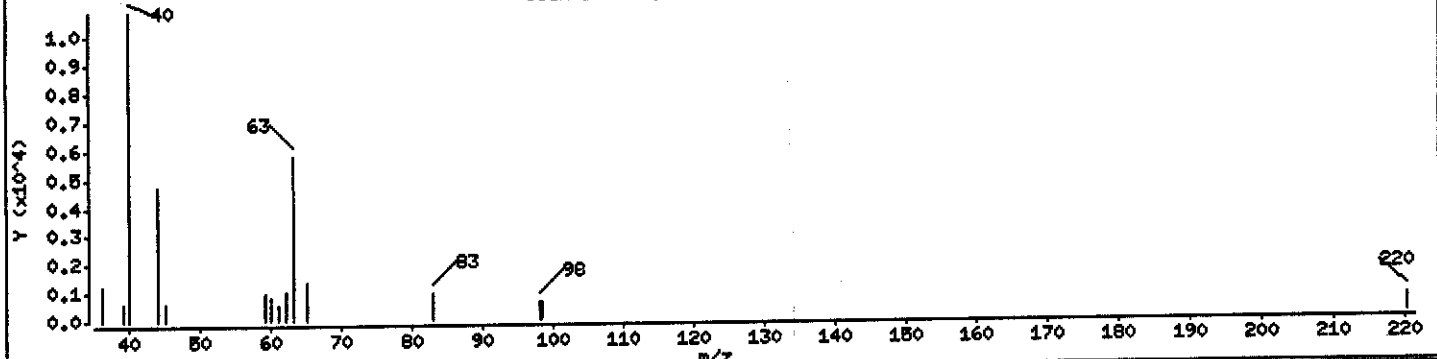
Column phase: DB624

Column diameter: 0.16

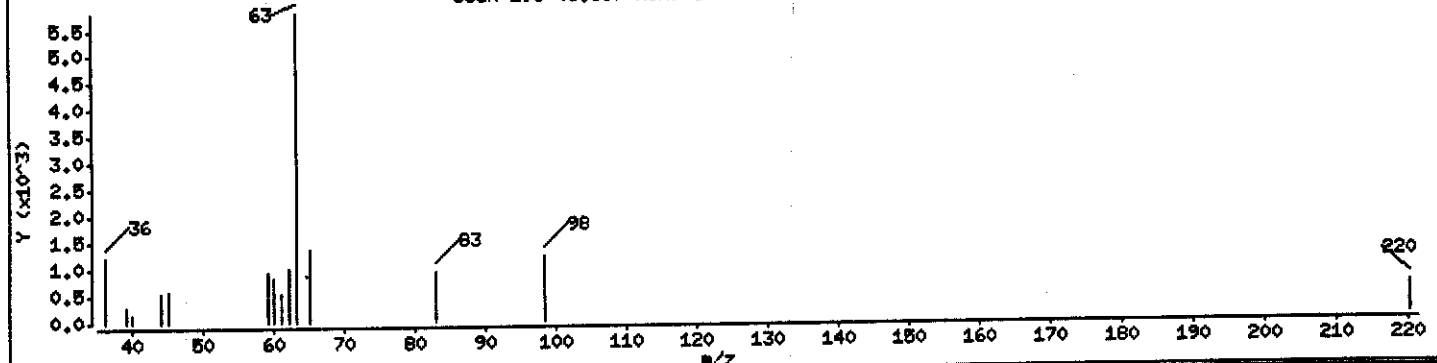
28 1,1-Dichloroethane

Concentration: 0.1653 ug/L

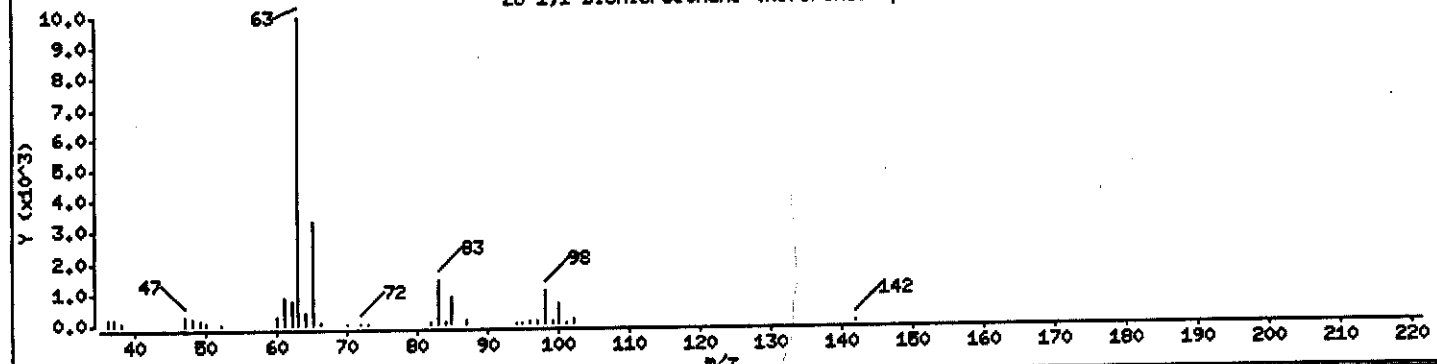
Scan 193 (3.657 min) of UXJ24287.D



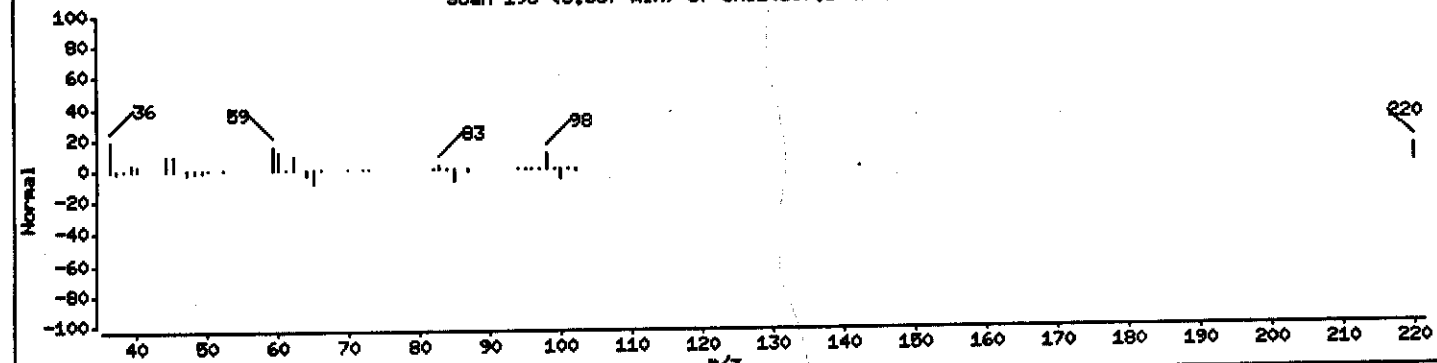
Scan 193 (3.657 min) of UXJ24287.D (Subtracted)



28 1,1-Dichloroethane (Reference Spectrum)



Scan 193 (3.657 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanch04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE840/35.5-40.5/092

Instrument: s3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

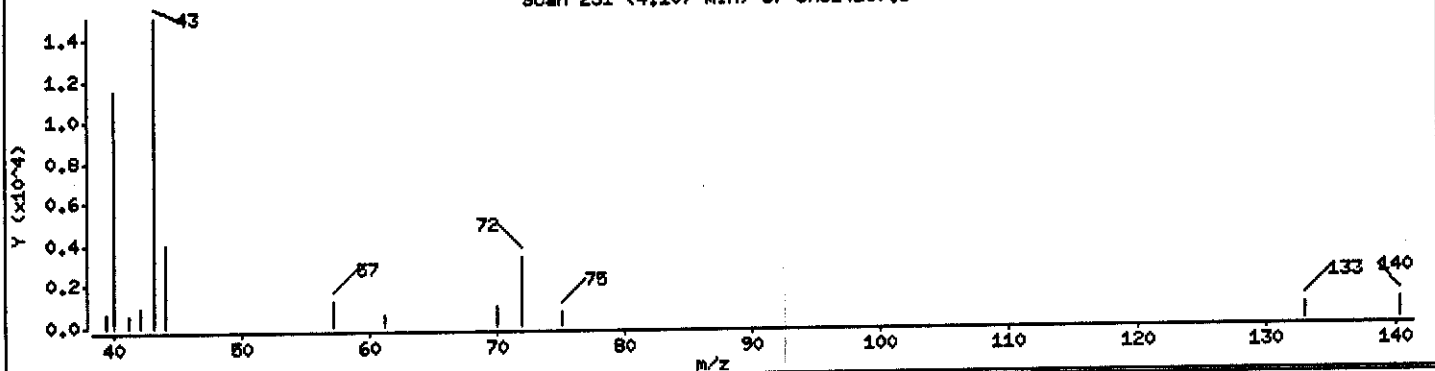
Column phase: DB624

Column diameter: 0.18

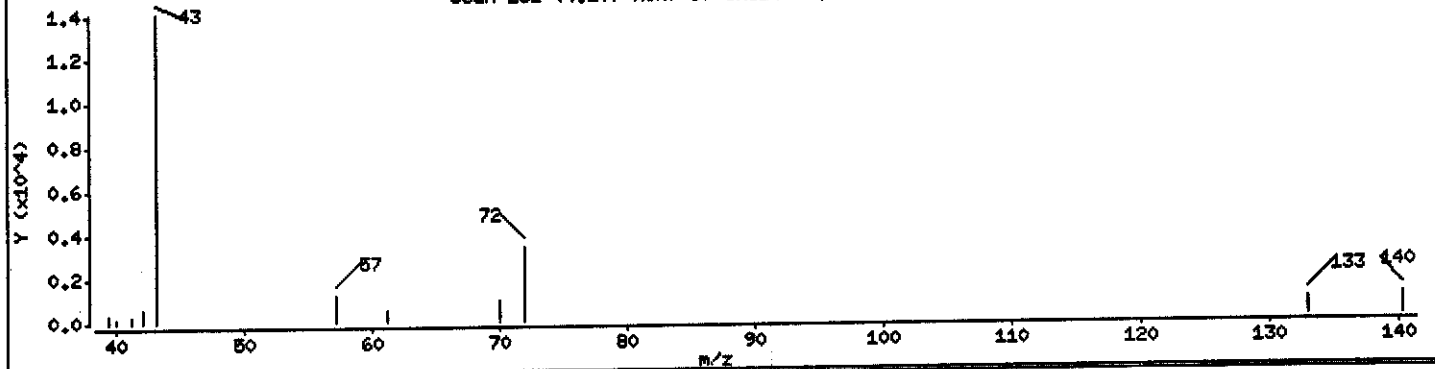
30 2-Butanone

Concentration: 1.652 ug/L

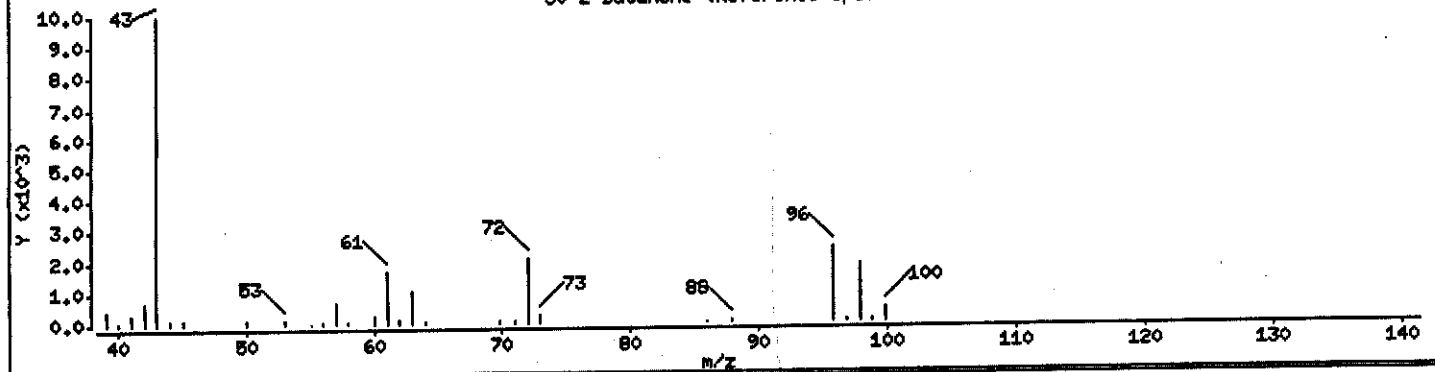
Scan 231 (4.107 min) of UXJ24287.D



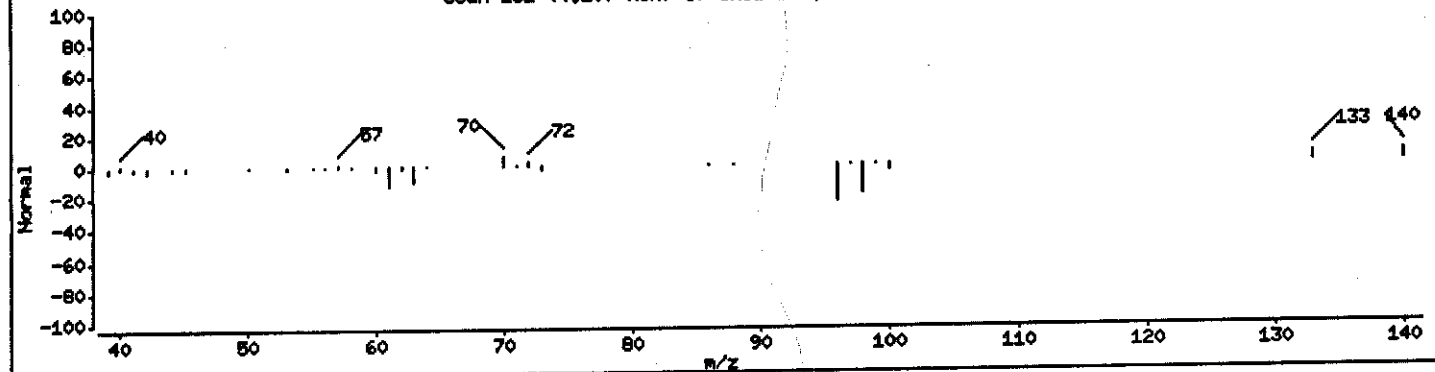
Scan 231 (4.107 min) of UXJ24287.D (Subtracted)



30 2-Butanone (Reference Spectrum)



Scan 231 (4.107 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VEB40/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

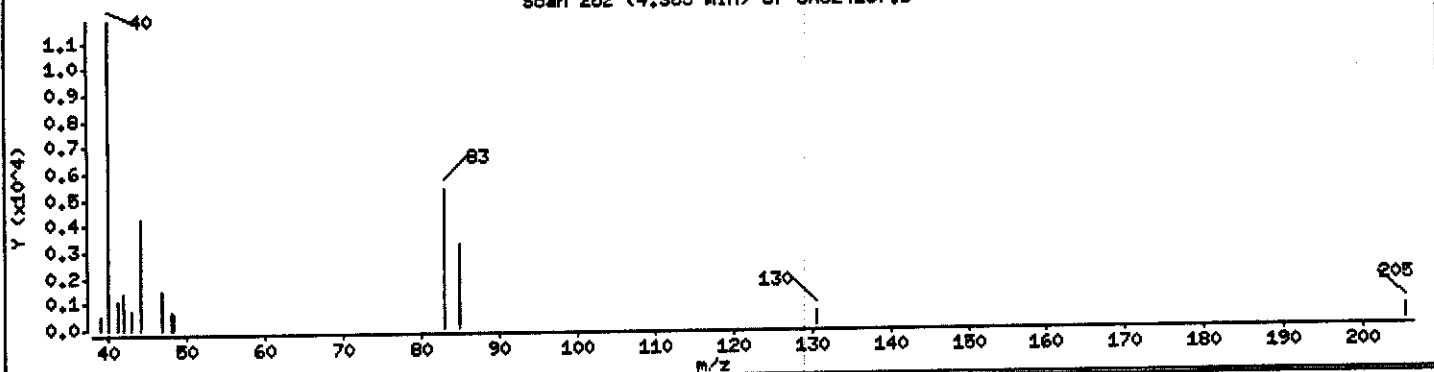
Column diameter: 0.18

Column phase: DB624

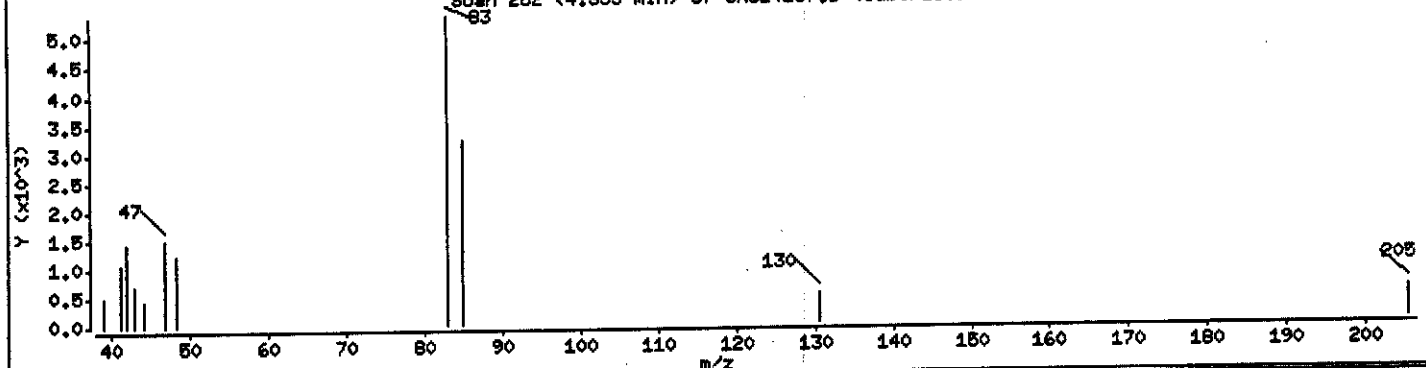
Concentration: 0.1754 ug/L

35 Chloroform

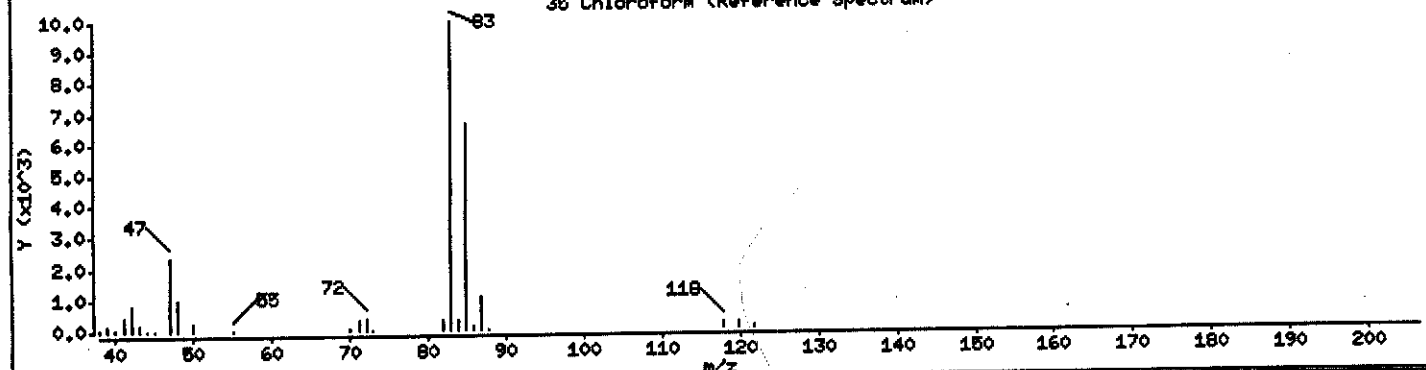
Scan 252 (4.355 min) of UXJ24287.D



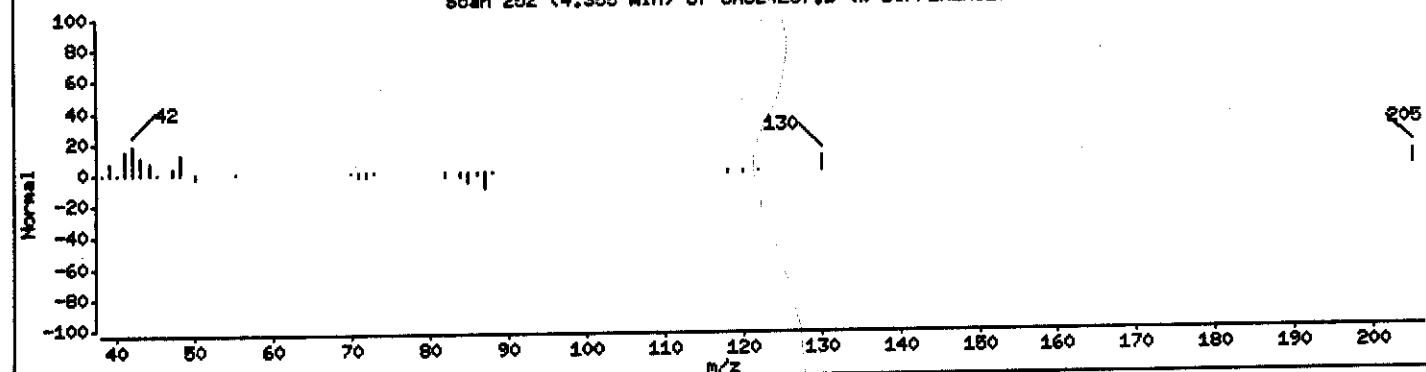
Scan 252 (4.355 min) of UXJ24287.D (Subtracted)



35 Chloroform (Reference Spectrum)



Scan 252 (4.355 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanch04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: CRDX11AA,5ML/5ML

Purge Volume: 5.0

Operator: 43882

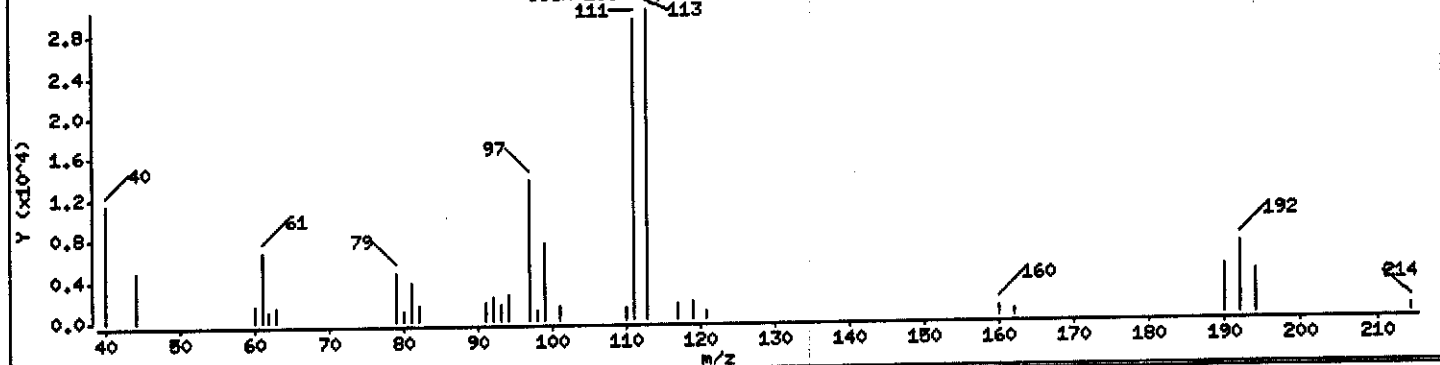
Column phase: DB624

Column diameter: 0.16

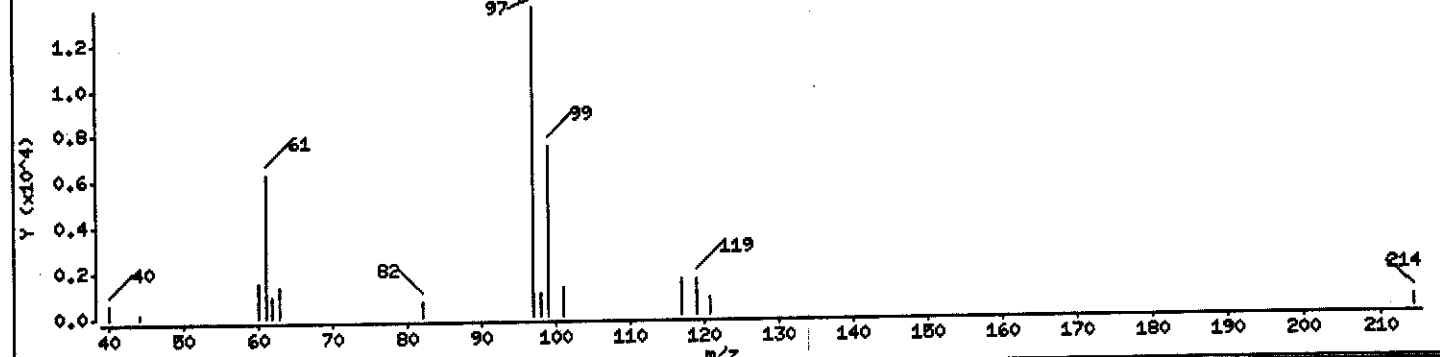
37 1,1,1-Trichloroethane

Concentration: 0.4768 ug/L

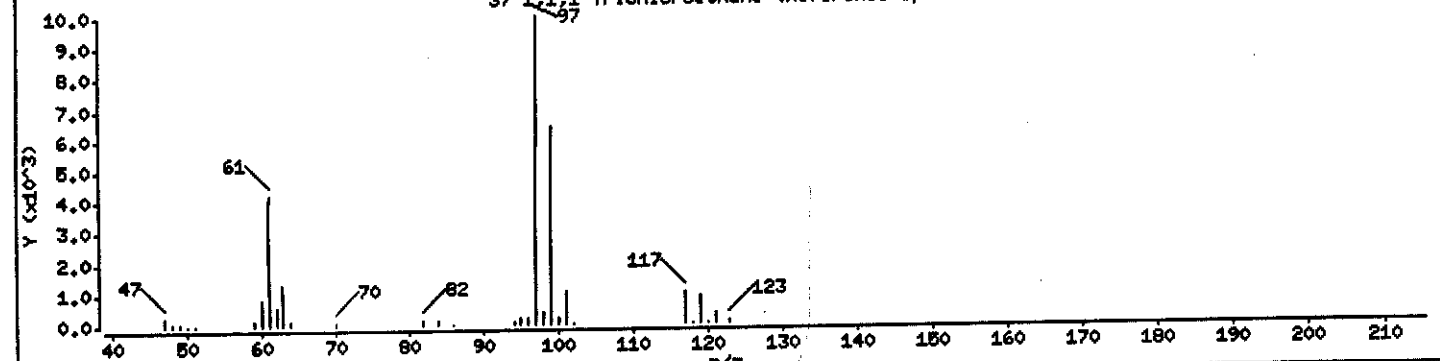
Scan 266 (4.521 min) of UXJ24287.D



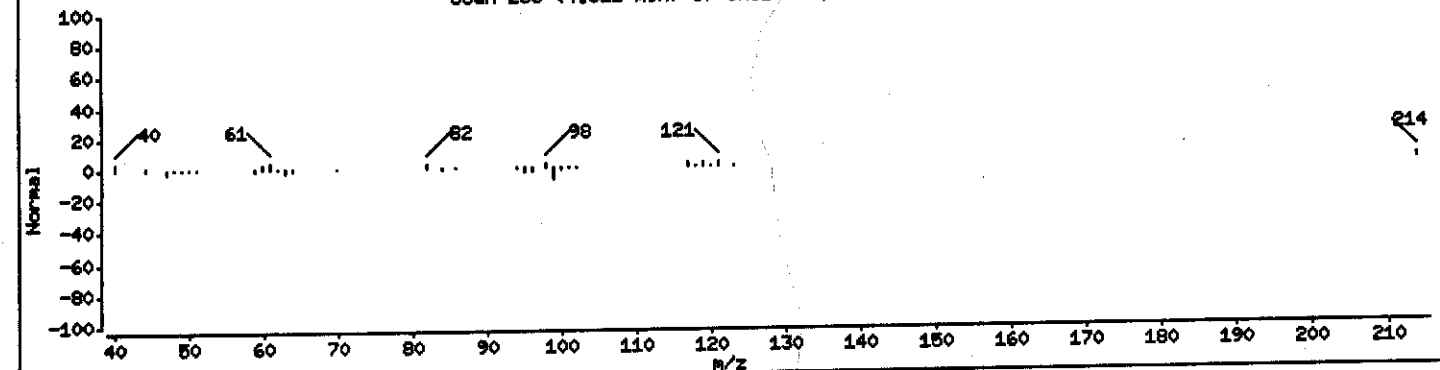
Scan 266 (4.521 min) of UXJ24287.D (Subtracted)



37 1,1,1-Trichloroethane (Reference Spectrum)



Scan 266 (4.521 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcaroh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: CRDX11AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

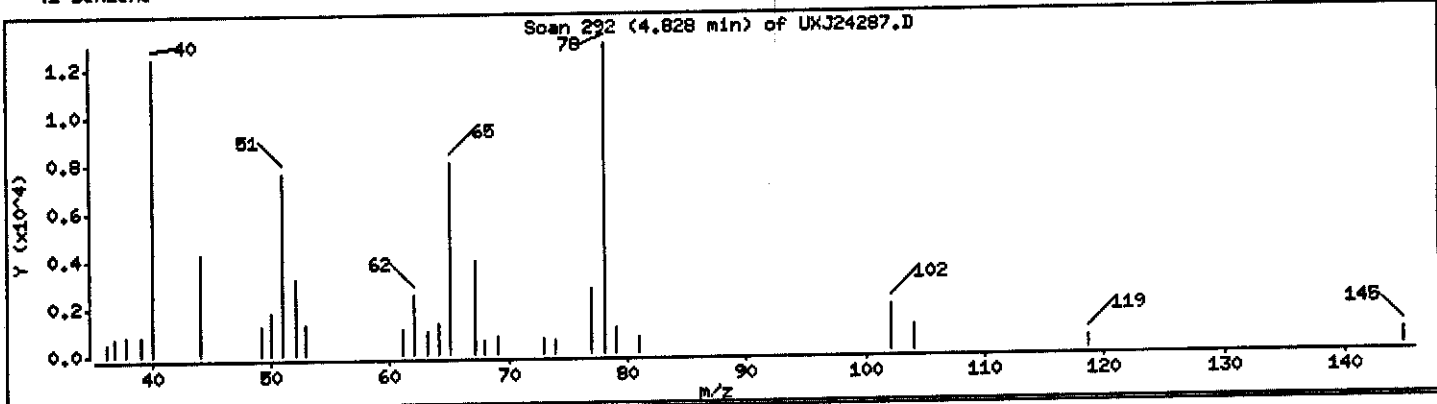
Column phase: DB624

Column diameter: 0.18

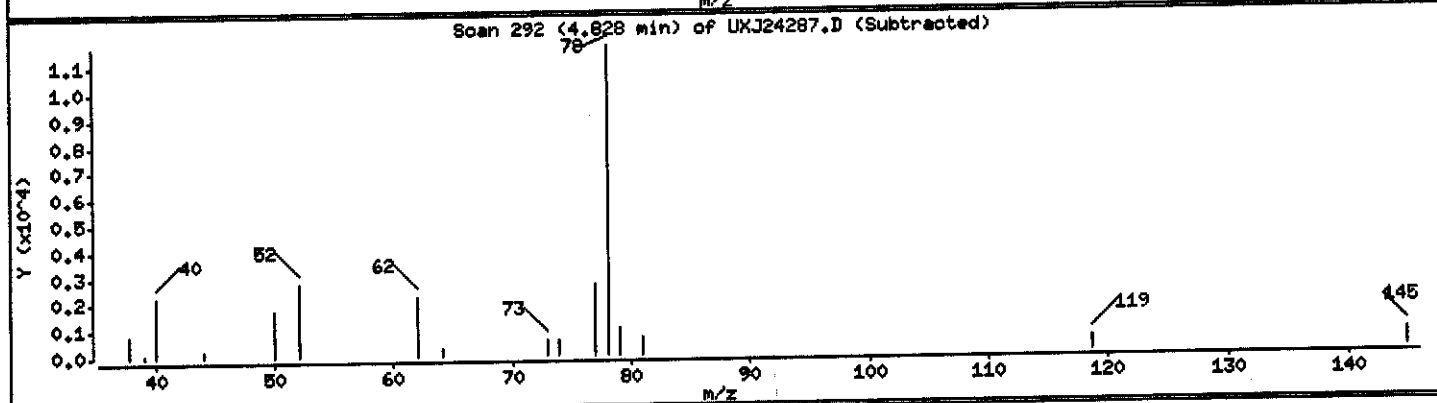
Concentration: 0.1711 ug/L

41 Benzene

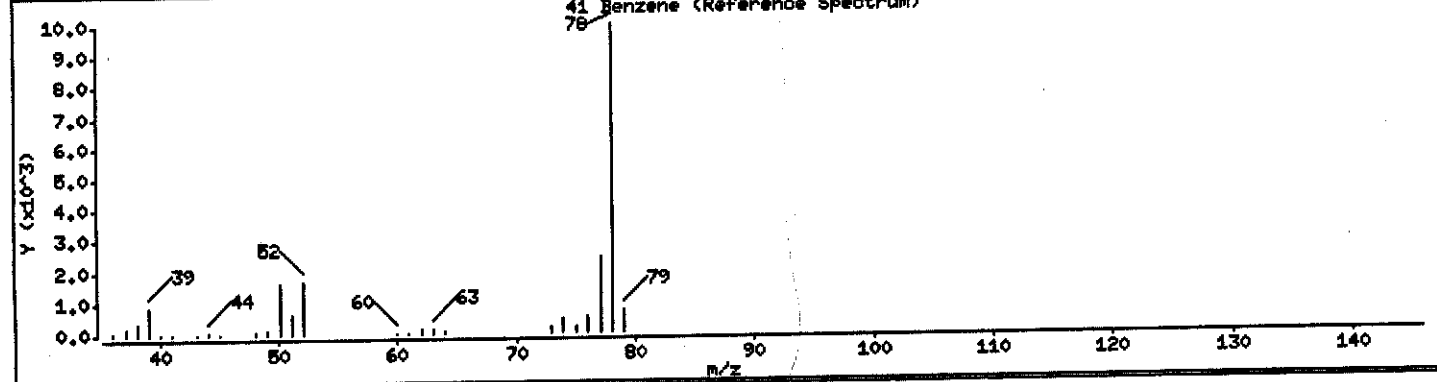
Scan 292 (4.828 min) of UXJ24287.D



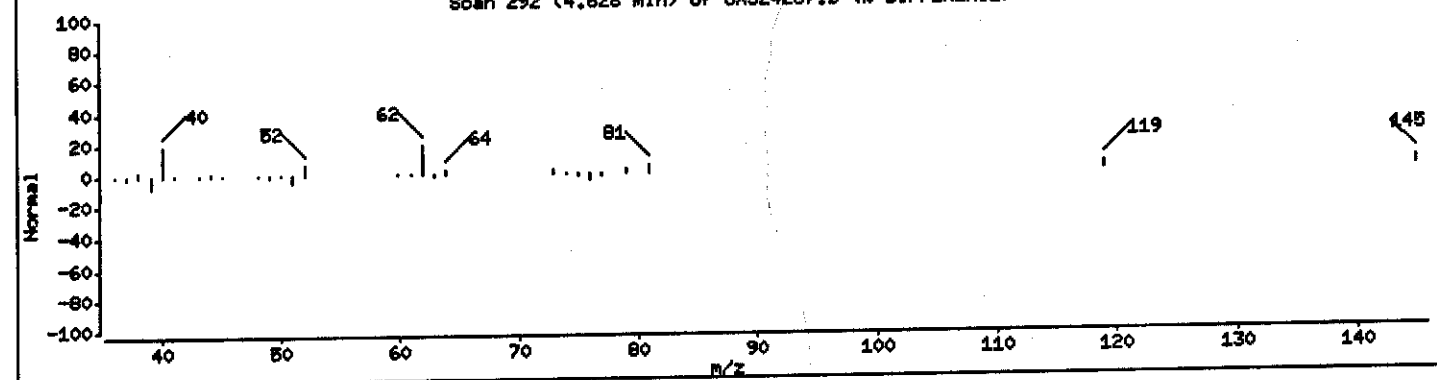
Scan 292 (4.828 min) of UXJ24287.D (Subtracted)



41 Benzene (Reference Spectrum)



Scan 292 (4.828 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

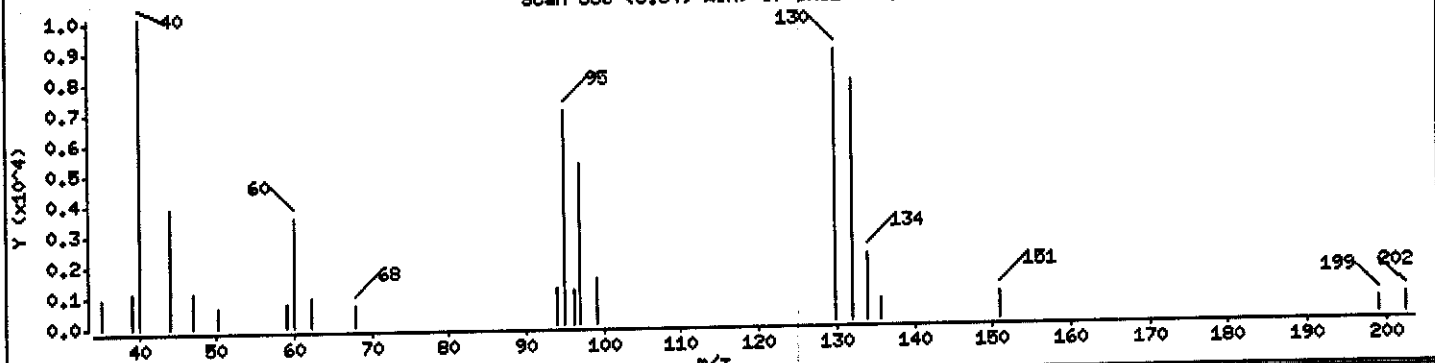
Column diameter: 0.18

Column phase: DB624

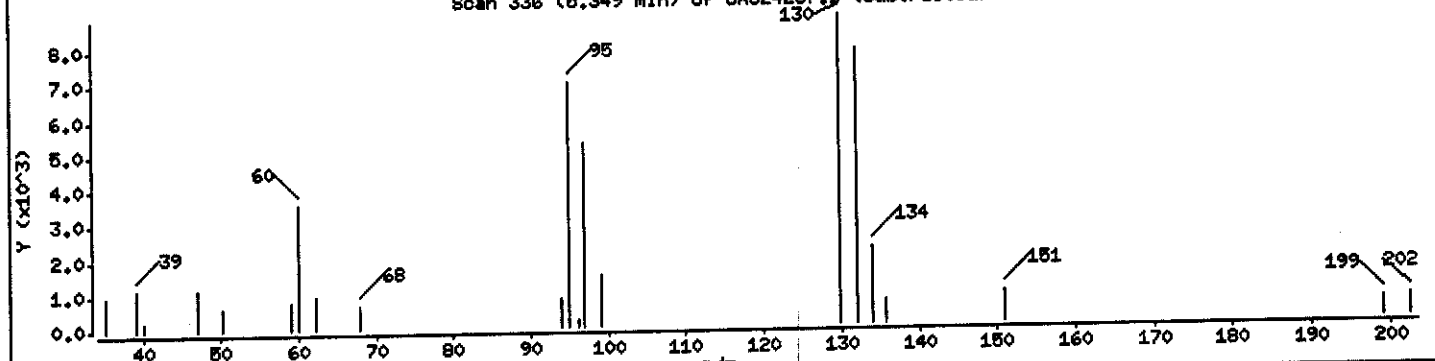
Concentration: 0.3903 ug/L

42 Trichloroethene

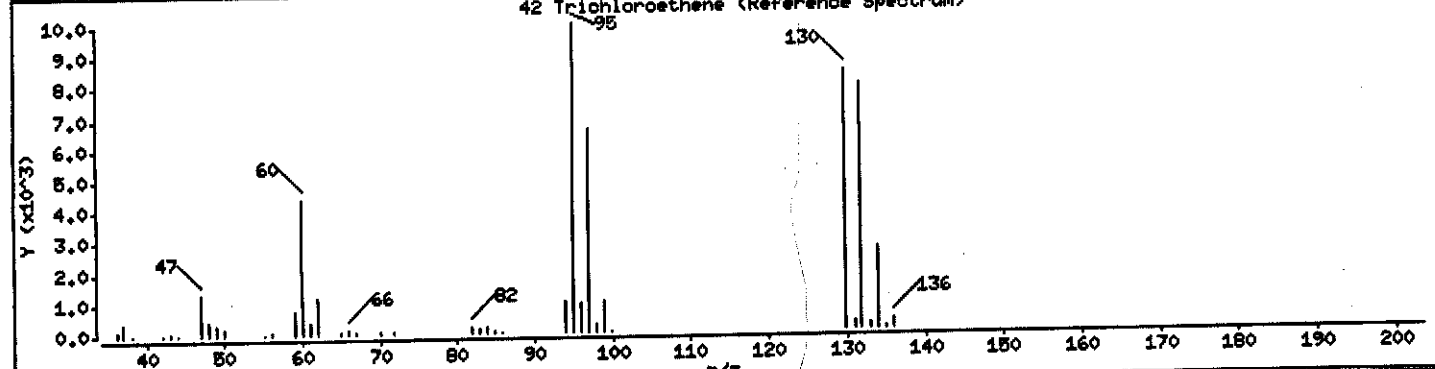
Scan 336 (5.349 min) of UXJ24287.D



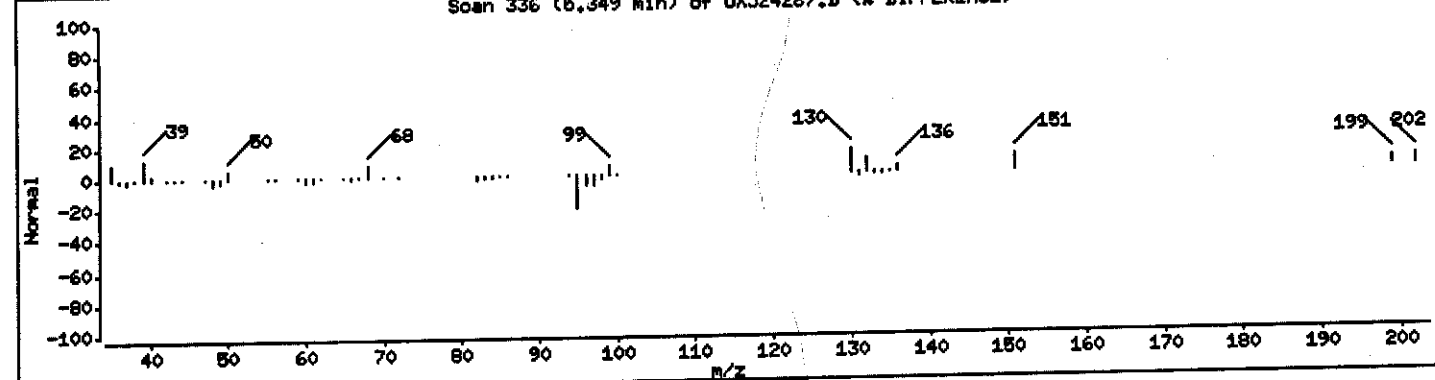
Scan 336 (5.349 min) of UXJ24287.D (Subtracted)



42 Trichloroethene (Reference Spectrum)



Scan 336 (5.349 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\HSV\z3ux11.i\J41001A.b\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE340/35.5-40.5/092

Instrument: z3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

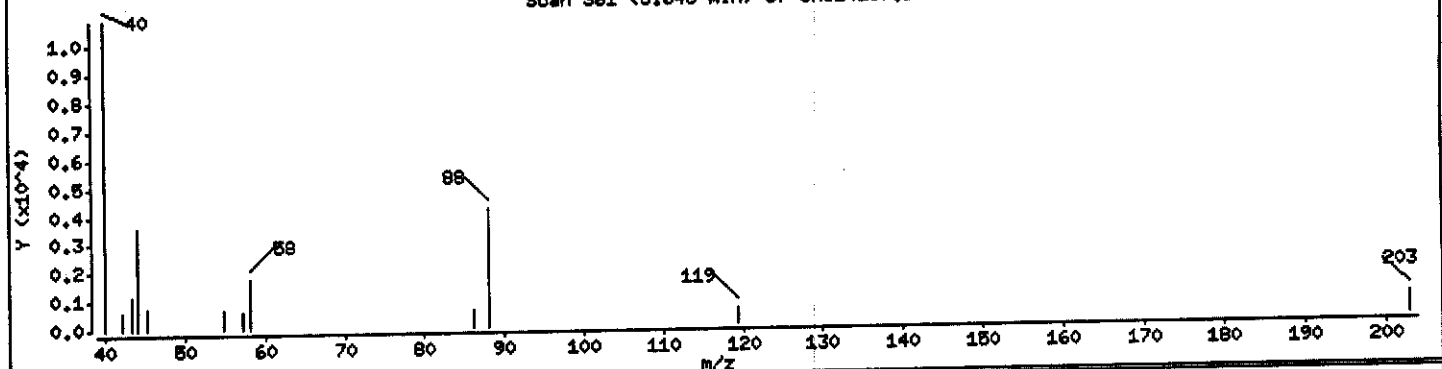
Column phase: DB624

Column diameter: 0.18

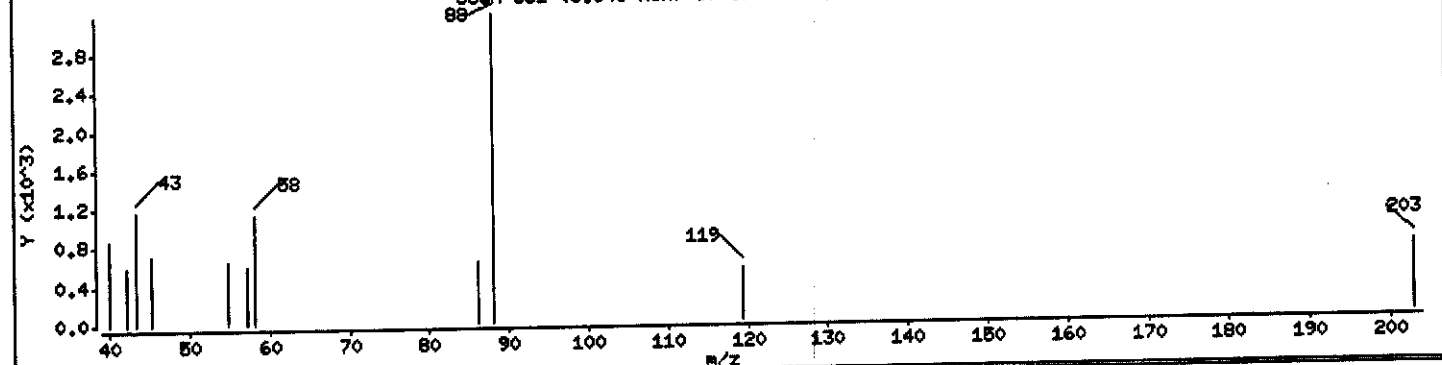
44 1,4-Dioxane

Concentration: 19.950 ug/L

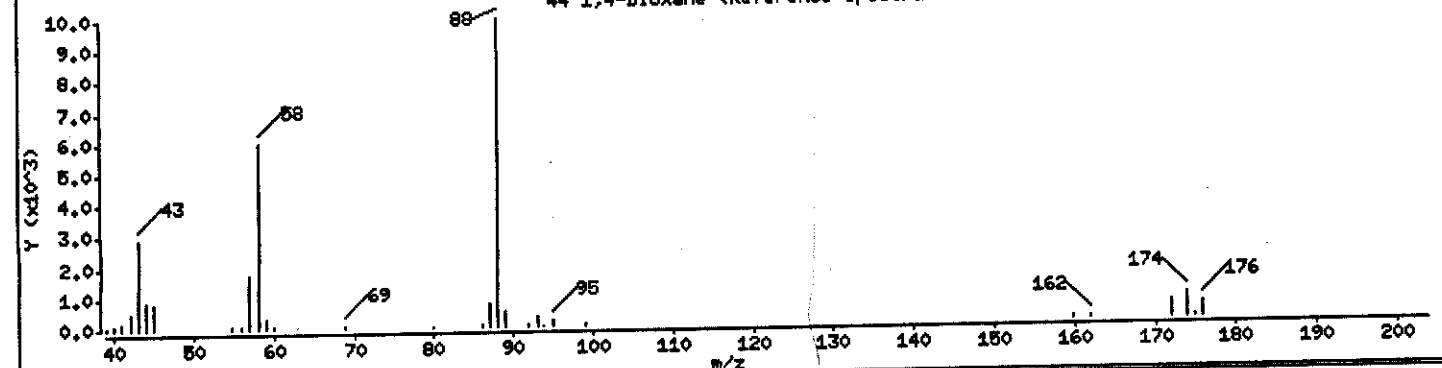
Scan 361 (5.645 min) of UXJ24287.D



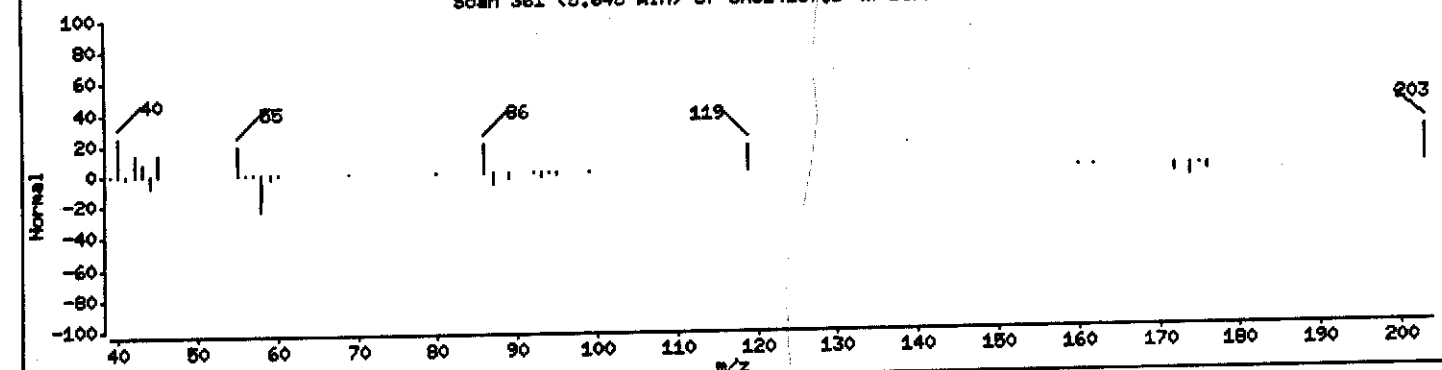
Scan 361 (5.645 min) of UXJ24287.D (Subtracted)



44 1,4-Dioxane (Reference Spectrum)



Scan 361 (5.645 min) of UXJ24287.D (% DIFFERENCE)



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J41001A.b\\UXJ24287.D

Date : 01-OCT-2004 11:49

Client ID: VE540/35.5-40.5/092

Instrument: a3ux11.i

Sample Info: GRDX11AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

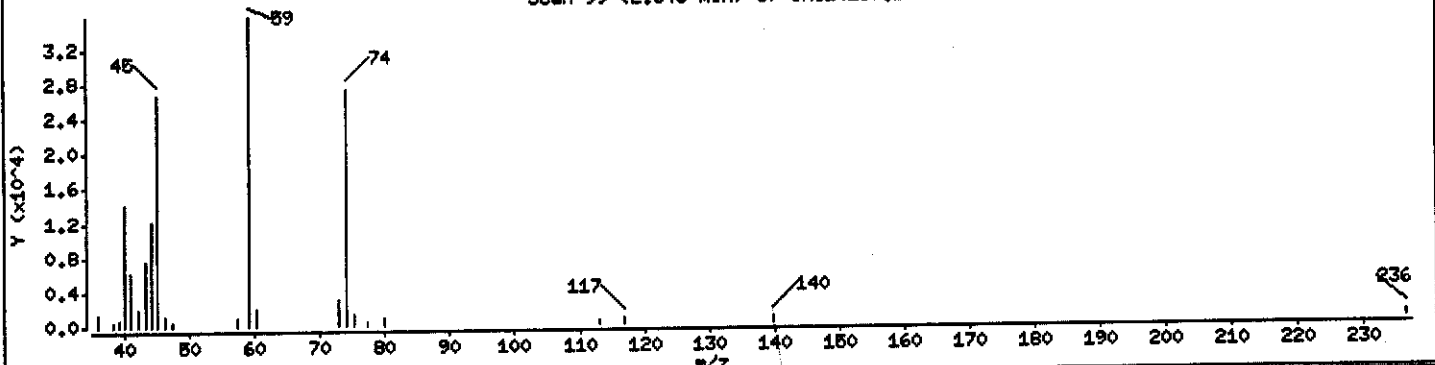
Column diameter: 0.18

Column phase: DB624

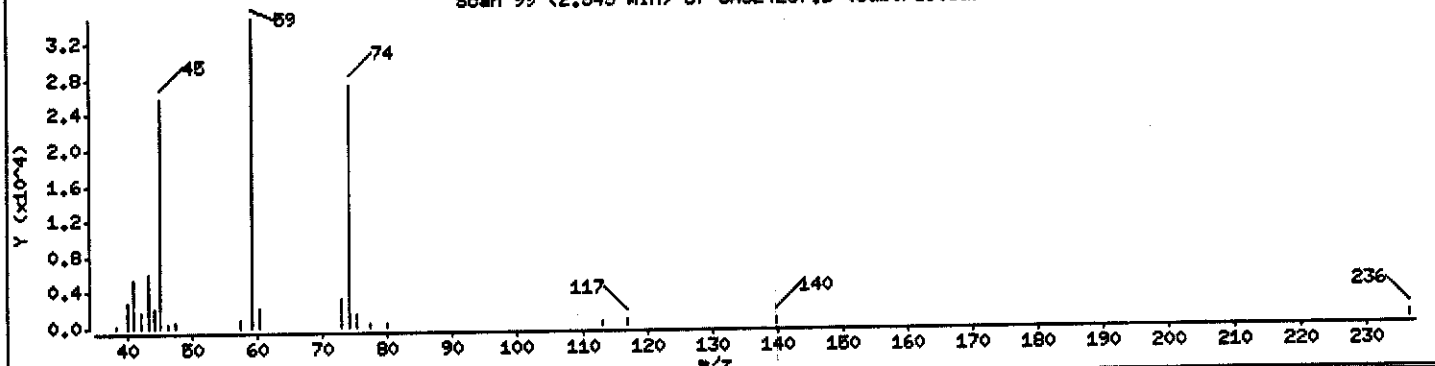
Concentration: 2.321 ug/L

89 Ethyl Ether

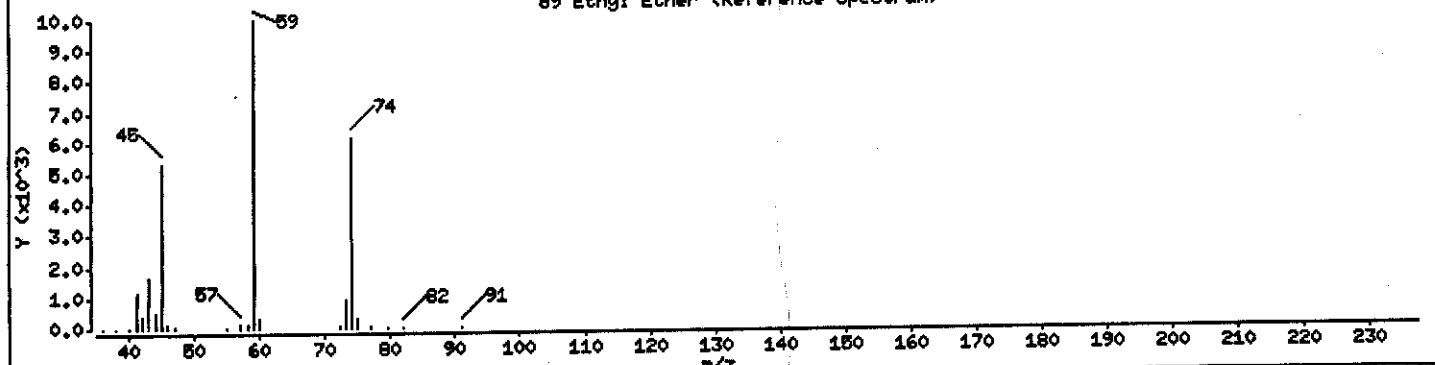
Scan 99 (2.545 min) of UXJ24287.D



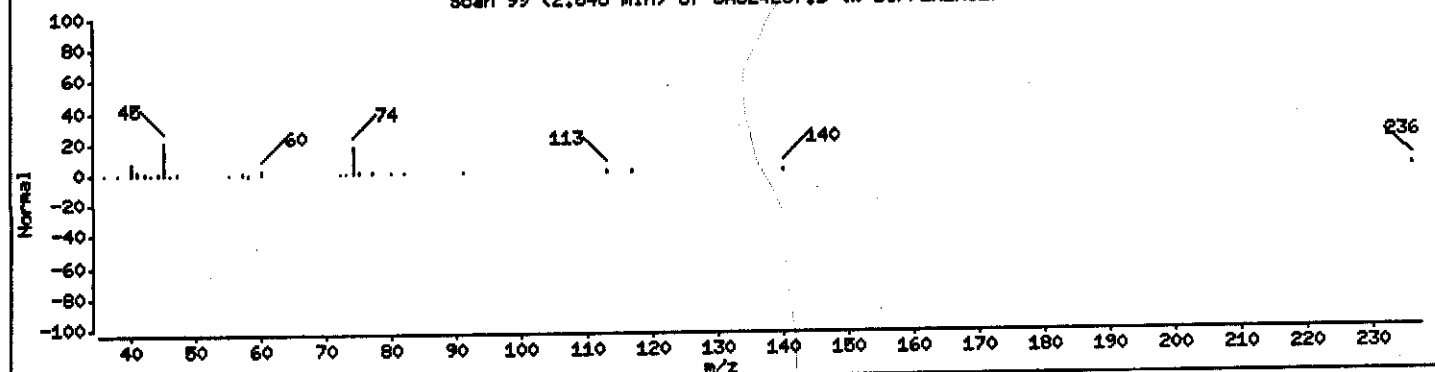
Scan 99 (2.545 min) of UXJ24287.D (Subtracted)



89 Ethyl Ether (Reference Spectrum)



Scan 99 (2.545 min) of UXJ24287.D (% DIFFERENCE)



PAYNE FIRM INC.

Client Sample ID: VE541/26.5-31.5/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-003 Work Order #....: GRDX61AA Matrix.....: WG
 Date Sampled....: 09/28/04 13:55 Date Received...: 09/29/04
 Prep Date.....: 10/01/04 Analysis Date...: 10/01/04
 Prep Batch #....: 4275213
 Dilution Factor: 1 Initial Wgt/Vol: 5 mL Final Wgt/Vol...: 5 mL
 Method.....: SW846 8260B

| PARAMETER | RESULT | REPORTING LIMIT | UNITS |
|------------------------------------|----------|-----------------|-------|
| Acetone | 5.3 J | 10 | ug/L |
| Acetonitrile | ND | 20 | ug/L |
| Acrolein | ND | 20 | ug/L |
| Acrylonitrile | ND | 20 | ug/L |
| Benzene | 0.24 J | 1.0 | ug/L |
| Bromodichloromethane | ND | 1.0 | ug/L |
| Bromoform | ND | 1.0 | ug/L |
| Bromomethane | ND | 1.0 | ug/L |
| 2-Butanone | 2.2 J | 10 | ug/L |
| Carbon disulfide | ND | 1.0 | ug/L |
| Carbon tetrachloride | ND | 1.0 | ug/L |
| Chlorobenzene | ND | 1.0 | ug/L |
| Chloroprene | ND | 2.0 | ug/L |
| Dibromochloromethane | ND | 1.0 | ug/L |
| Chloroethane | ND | 1.0 | ug/L |
| Chloroform | 0.27 J | 1.0 | ug/L |
| Chloromethane | 0.29 J,B | 1.0 | ug/L |
| 3-Chloropropene | ND | 2.0 | ug/L |
| 1,2-Dibromo-3-chloropropane (DBCP) | ND | 2.0 | ug/L |
| 1,2-Dibromoethane | ND | 1.0 | ug/L |
| Dibromomethane | ND | 1.0 | ug/L |
| trans-1,4-Dichloro-2-butene | ND | 1.0 | ug/L |
| 1,1-Dichloroethane | ND | 1.0 | ug/L |
| 1,2-Dichloroethane | 2.4 | 1.0 | ug/L |
| cis-1,2-Dichloroethene | 0.90 J | 1.0 | ug/L |
| trans-1,2-Dichloroethene | ND | 1.0 | ug/L |
| 1,1-Dichloroethene | ND | 1.0 | ug/L |
| 1,2-Dichloroethene (total) | 0.90 J | 2.0 | ug/L |
| Dichlorofluoromethane | ND | 2.0 | ug/L |
| 1,2-Dichloropropane | ND | 1.0 | ug/L |
| cis-1,3-Dichloropropene | ND | 1.0 | ug/L |
| trans-1,3-Dichloropropene | ND | 1.0 | ug/L |
| 1,4-Dioxane | 18 J | 50 | ug/L |
| Ethylbenzene | ND | 1.0 | ug/L |
| Ethyl methacrylate | ND | 1.0 | ug/L |

(Continued on next page)

PAYNE FIRM INC.

Client Sample ID: VE541/26.5-31.5/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-003 Work Order #....: GRDX61AA Matrix.....: WG

| <u>PARAMETER</u> | <u>RESULT</u> | <u>REPORTING</u> | |
|---------------------------|---------------|------------------|--------------|
| | | <u>LIMIT</u> | <u>UNITS</u> |
| 2-Hexanone | ND | 10 | ug/L |
| Iodomethane | ND | 1.0 | ug/L |
| Isobutanol | ND | 50 | ug/L |
| Methacrylonitrile | ND | 2.0 | ug/L |
| Methylene chloride | ND | 1.0 | ug/L |
| Methyl methacrylate | ND | 2.0 | ug/L |
| 4-Methyl-2-pentanone | ND | 10 | ug/L |
| Propionitrile | ND | 4.0 | ug/L |
| Styrene | ND | 1.0 | ug/L |
| 1,1,1,2-Tetrachloroethane | ND | 1.0 | ug/L |
| 1,1,2,2-Tetrachloroethane | ND | 1.0 | ug/L |
| Tetrachloroethene | ND | 1.0 | ug/L |
| Toluene | ND | 1.0 | ug/L |
| 1,1,1-Trichloroethane | ND | 1.0 | ug/L |
| 1,1,2-Trichloroethane | ND | 1.0 | ug/L |
| Trichloroethene | 0.38 J | 1.0 | ug/L |
| Trichlorofluoromethane | ND | 1.0 | ug/L |
| 1,2,3-Trichloropropane | ND | 1.0 | ug/L |
| Vinyl acetate | ND | 2.0 | ug/L |
| Vinyl chloride | ND | 1.0 | ug/L |
| Xylenes (total) | ND | 2.0 | ug/L |

| <u>SURROGATE</u> | <u>PERCENT RECOVERY</u> | <u>RECOVERY</u> |
|-----------------------|-----------------------------|-----------------|
| | | <u>LIMITS</u> |
| Dibromofluoromethane | 112 | (73 - 122) |
| 1,2-Dichloroethane-d4 | 113 | (61 - 128) |
| Toluene-d8 | 86 | (76 - 110) |
| 4-Bromofluorobenzene | 81 | (74 - 116) |

NOTE (S) :

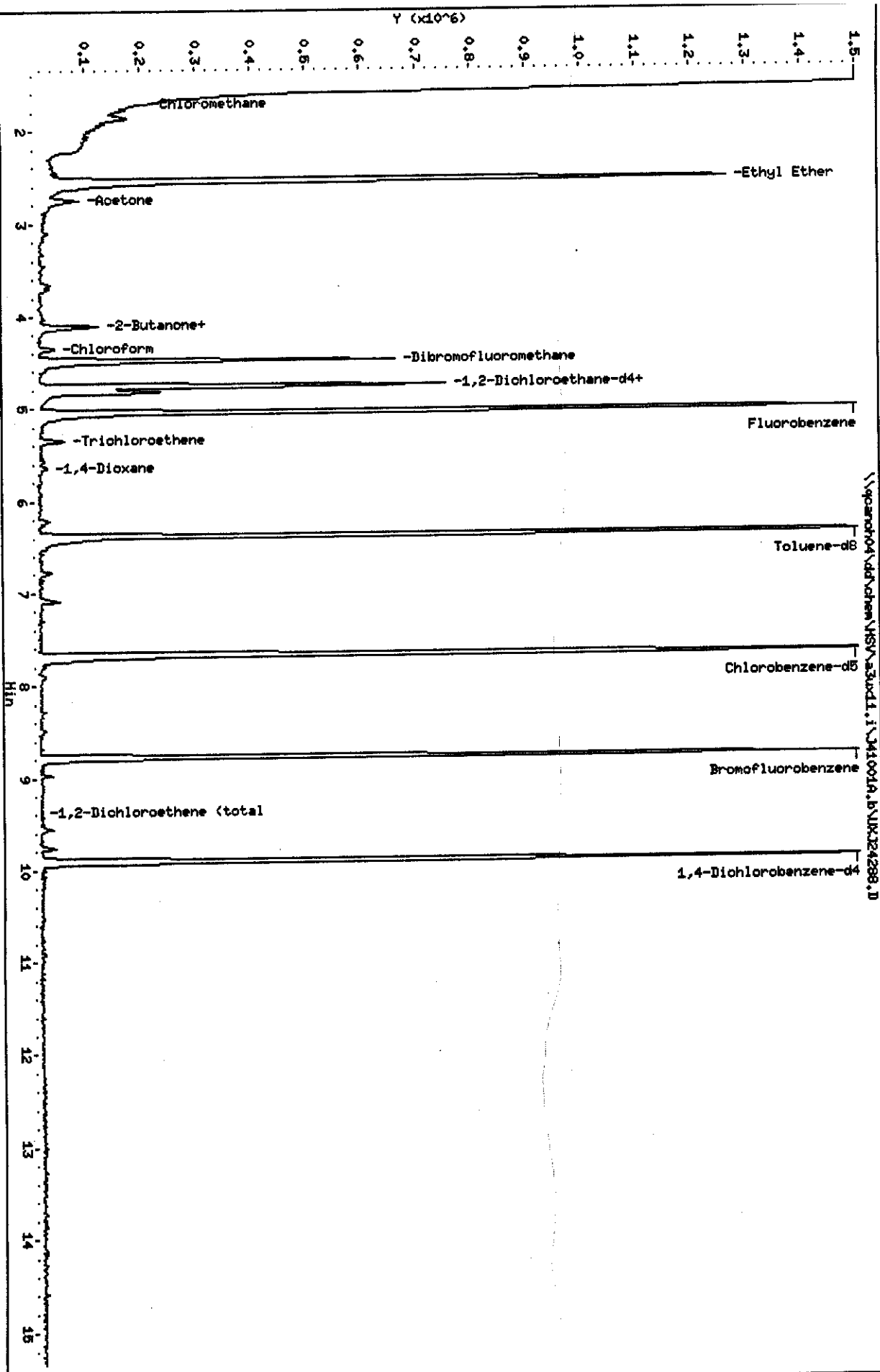
J Estimated result. Result is less than RL.

B Method blank contamination. The associated method blank contains the target analyte at a reportable level.

Data File: \\pcanon04\\dat\\chem\\HSV\\a30x1.i\\J41001A.b\\UX24288.D
Date : 01-OCT-2004 12:28
Client ID: VEB41/26.5-31.5/092
Sample Info: GRDX6100,5HL/5HL
Purge Volume: 5.0

Column phase: DB624

Instrument: z30x1.i
Operator: 43682
Column diameter: 0.48



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24288.D
Report Date: 04-Oct-2004 10:01

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24288.D
Lab Smp Id: GRDX61AA Client Smp ID: VE541/26.5-31.5/092
Inj Date : 01-OCT-2004 12:28 Inst ID: a3ux11.i
Operator : 43582
Smp Info : GRDX61AA, 5ML/5ML
Misc Info : J41001A, 8260LLUX11,, 43582
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 12
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 4-8260+IX.sub
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | |
|----------------------------|-----------|------------------------|---------------|---------|---------|----------|-------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) |
| * 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 1815392 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1560887 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 683725 | 50.0000 | | |
| \$ 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 471305 | 55.7608 | 11.152 | |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 655941 | 56.4371 | 11.287 | |
| \$ 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 1621664 | 43.2459 | 8.649 | |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 643041 | 40.3594 | 8.072 | |
| 8 Dichlorodifluoromethane | 85 | Compound Not Detected. | | | | | |
| 9 Chloromethane | 50 | 1.704 | 1.704 (0.338) | 24947 | 1.47219 | 0.2944 | |
| 10 Vinyl Chloride | 62 | Compound Not Detected. | | | | | |
| 11 Bromomethane | 94 | Compound Not Detected. | | | | | |
| 12 Chloroethane | 64 | Compound Not Detected. | | | | | |
| 13 Trichlorofluoromethane | 101 | Compound Not Detected. | | | | | |
| 15 Acrolein | 56 | Compound Not Detected. | | | | | |
| 16 Acetone | 43 | 2.745 | 2.745 (0.545) | 123946 | 26.5666 | 5.313 | |
| 17 1,1-Dichloroethene | 96 | Compound Not Detected. | | | | | |
| 18 Freon-113 | 151 | Compound Not Detected. | | | | | |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | (ng) | (ug/L) |
|---------------------------------|-----------|------|----------------|---------------|--------|------------------------|---------|---------|
| | | | RT | EXP RT | REL RT | RESPONSE | | |
| 19 Iodomethane | | 142 | | | | Compound Not Detected. | | |
| 20 Carbon Disulfide | | 76 | | | | Compound Not Detected. | | |
| 21 Methylene Chloride | | 84 | | | | Compound Not Detected. | | |
| 22 Acetonitrile | | 41 | | | | Compound Not Detected. | | |
| 23 Acrylonitrile | | 53 | | | | Compound Not Detected. | | |
| 24 Methyl tert-butyl ether | | 73 | | | | Compound Not Detected. | | |
| 25 trans-1,2-Dichloroethene | | 96 | | | | Compound Not Detected. | | |
| 26 Hexane | | 86 | | | | Compound Not Detected. | | |
| 27 Vinyl acetate | | 43 | | | | Compound Not Detected. | | |
| 28 1,1-Dichloroethane | | 63 | | | | Compound Not Detected. | | |
| 29 tert-Butyl Alcohol | | 59 | | | | Compound Not Detected. | | |
| 30 2-Butanone | | 43 | 4.106 | 4.106 (0.815) | | 53695 | 10.7987 | 2.160 |
| M 31 1,2-Dichloroethene (total) | | 96 | | | | 45528 | 4.49920 | 0.8998 |
| 32 cis-1,2-dichloroethene | | 96 | 4.106 | 4.106 (0.815) | | 45528 | 4.49920 | 0.8998 |
| 33 2,2-Dichloropropane | | 77 | | | | Compound Not Detected. | | |
| 34 Bromochloromethane | | 128 | | | | Compound Not Detected. | | |
| 35 Chloroform | | 83 | 4.355 | 4.355 (0.864) | | 23439 | 1.32714 | 0.2654 |
| 36 Tetrahydrofuran | | 42 | | | | Compound Not Detected. | | |
| 37 1,1,1-Trichloroethane | | 97 | | | | Compound Not Detected. | | |
| 38 1,1-Dichloropropene | | 75 | | | | Compound Not Detected. | | |
| 39 Carbon Tetrachloride | | 117 | | | | Compound Not Detected. | | |
| 40 1,2-Dichloroethane | | 62 | 4.816 | 4.816 (0.955) | | 171727 | 11.9004 | 2.380 |
| 41 Benzene | | 78 | 4.828 | 4.828 (0.958) | | 49462 | 1.18341 | 0.2367 |
| 42 Trichloroethene | | 130 | 5.349 | 5.349 (1.061) | | 18027 | 1.88984 | 0.3780 |
| 43 1,2-Dichloropropene | | 63 | | | | Compound Not Detected. | | |
| 44 1,4-Dioxane | | 88 | 5.633 | 5.633 (1.117) | | 8071 | 91.3251 | 18.265 |
| 45 Dibromomethane | | 93 | | | | Compound Not Detected. | | |
| 46 Bromodichloromethane | | 83 | | | | Compound Not Detected. | | |
| 47 2-Chloroethyl vinyl ether | | 63 | | | | Compound Not Detected. | | |
| 48 cis-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | | |
| 49 4-Methyl-2-pentanone | | 43 | | | | Compound Not Detected. | | |
| 50 Toluene | | 91 | | | | Compound Not Detected. | | |
| 51 trans-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | | |
| 52 Ethyl Methacrylate | | 69 | | | | Compound Not Detected. | | |
| 53 1,1,2-Trichloroethane | | 97 | | | | Compound Not Detected. | | |
| 54 1,3-Dichloropropane | | 76 | | | | Compound Not Detected. | | |
| 55 Tetrachloroethene | | 164 | | | | Compound Not Detected. | | |
| 56 2-Hexanone | | 43 | | | | Compound Not Detected. | | |
| 57 Dibromochloromethane | | 129 | | | | Compound Not Detected. | | |
| 58 1,2-Dibromoethane | | 107 | | | | Compound Not Detected. | | |
| 59 Chlorobenzene | | 112 | | | | Compound Not Detected. | | |
| 60 1,1,1,2-Tetrachloroethane | | 131 | | | | Compound Not Detected. | | |
| 61 Ethylbenzene | | 106 | | | | Compound Not Detected. | | |
| 62 m + p-Xylene | | 106 | | | | Compound Not Detected. | | |
| M 63 Xlenes (total) | | 106 | | | | Compound Not Detected. | | |
| 64 Xylene-o | | 106 | | | | Compound Not Detected. | | |
| 65 Styrene | | 104 | | | | Compound Not Detected. | | |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|--------------------------------|-----------|------|----------------|---------------|---------|------------------------|-------------------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 66 Bromoform | | 173 | | | | Compound Not Detected. | |
| 67 Isopropylbenzene | | 105 | | | | Compound Not Detected. | |
| 68 1,1,2,2-Tetrachloroethane | | 83 | | | | Compound Not Detected. | |
| 69 1,4-Dichloro-2-butene | | 53 | | | | Compound Not Detected. | |
| 70 1,2,3-Trichloropropane | | 110 | | | | Compound Not Detected. | |
| 71 Bromobenzene | | 156 | | | | Compound Not Detected. | |
| 72 n-Propylbenzene | | 120 | | | | Compound Not Detected. | |
| 73 2-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 74 1,3,5-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 75 4-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 76 tert-Butylbenzene | | 119 | | | | Compound Not Detected. | |
| 77 1,2,4-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 78 sec-Butylbenzene | | 105 | | | | Compound Not Detected. | |
| 79 4-Isopropyltoluene | | 119 | | | | Compound Not Detected. | |
| 80 1,3-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 81 1,4-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 82 n-Butylbenzene | | 91 | | | | Compound Not Detected. | |
| 83 1,2-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 84 1,2-Dibromo-3-chloropropane | | 157 | | | | Compound Not Detected. | |
| 85 1,2,4-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 86 Hexachlorobutadiene | | 225 | | | | Compound Not Detected. | |
| 87 Naphthalene | | 128 | | | | Compound Not Detected. | |
| 88 1,2,3-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 14 Dichlorofluoromethane | | 67 | | | | Compound Not Detected. | |
| 89 Ethyl Ether | | 59 | 2.544 | 2.532 (0.505) | 1156071 | 129.149 | 25.830 |
| 91 3-Chloropropene | | 76 | | | | Compound Not Detected. | |
| 92 Isopropyl Ether | | 87 | | | | Compound Not Detected. | |
| 93 2-Chloro-1,3-butadiene | | 53 | | | | Compound Not Detected. | |
| 94 Propionitrile | | 54 | | | | Compound Not Detected. | |
| 95 Ethyl Acetate | | 43 | | | | Compound Not Detected. | |
| 96 Methacrylonitrile | | 41 | | | | Compound Not Detected. | |
| 97 Isobutanol | | 41 | | | | Compound Not Detected. | |
| 99 n-Butanol | | 56 | | | | Compound Not Detected. | |
| 100 Methyl Methacrylate | | 41 | | | | Compound Not Detected. | |
| 101 2-Nitropropane | | 41 | | | | Compound Not Detected. | |
| 103 Cyclohexanone | | 55 | | | | Compound Not Detected. | |
| 98 Cyclohexane | | 56 | | | | Compound Not Detected. | |
| 143 Methyl Acetate | | 43 | | | | Compound Not Detected. | |
| 144 Methylcyclohexane | | 83 | | | | Compound Not Detected. | |
| 141 1,3,5-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |

Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VEB41/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: GRDX61AA,5ML/5ML

Purge Volume: 5.0

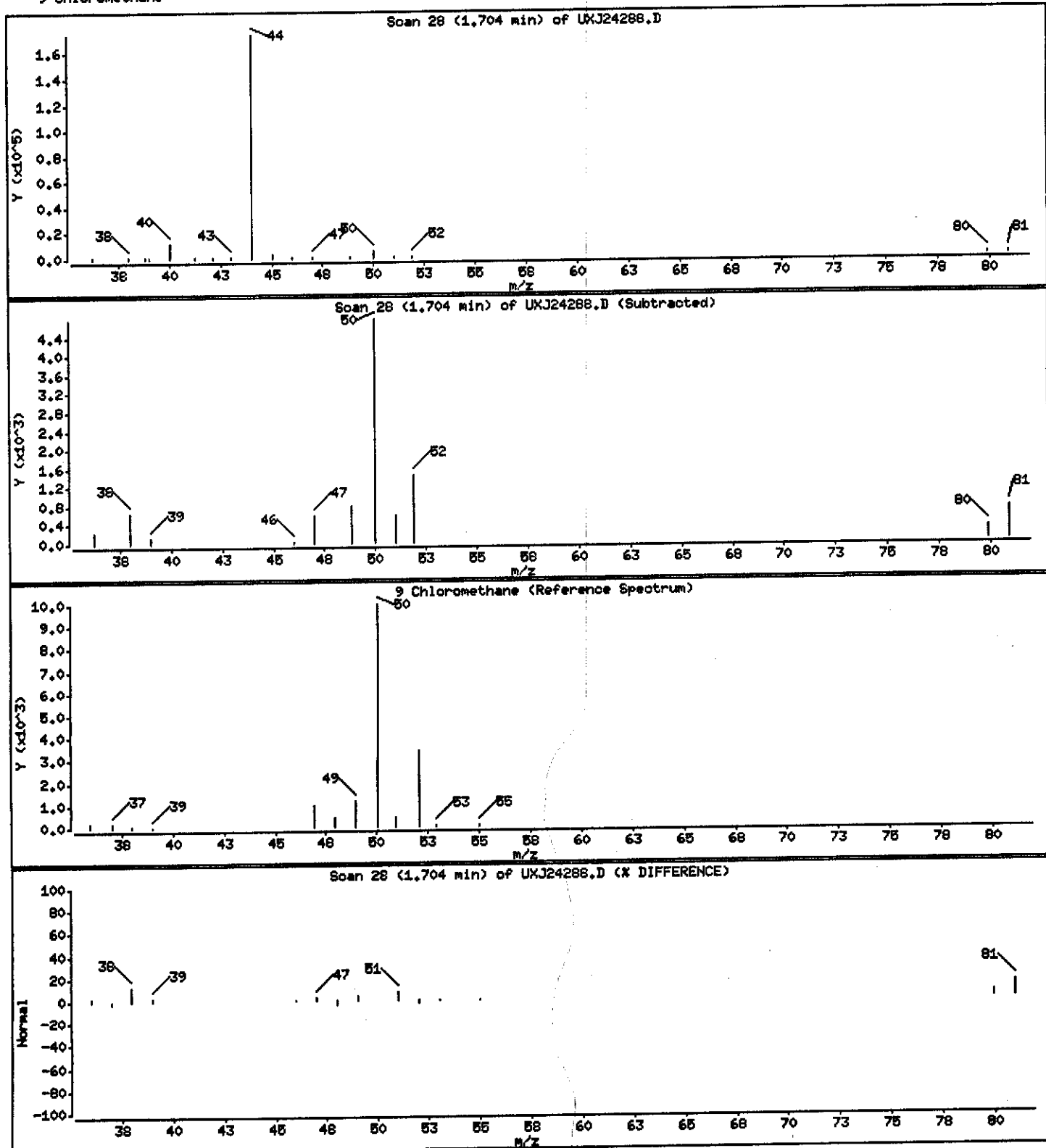
Operator: 43582

Column phase: DB624

Column diameter: 0.18

9 Chloromethane

Concentration: 0.2944 ug/L



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VE541/26.5-31.5/092

Sample Info: GRDX61AA,5ML/5ML

Purge Volume: 5.0

Column phase: DB624

Instrument: z3ux11.i

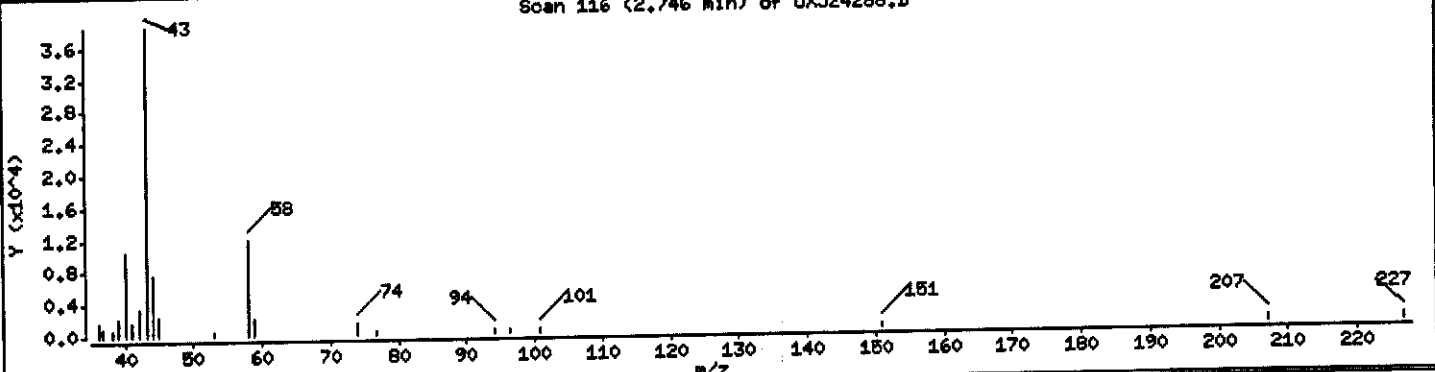
Operator: 43582

Column diameter: 0.18

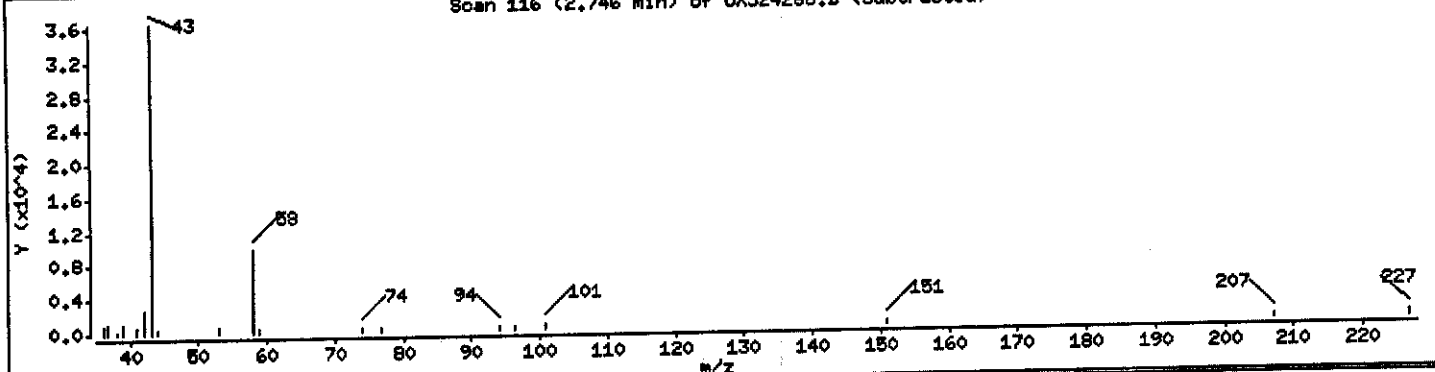
Concentration: 5.313 ug/L

16 Acetone

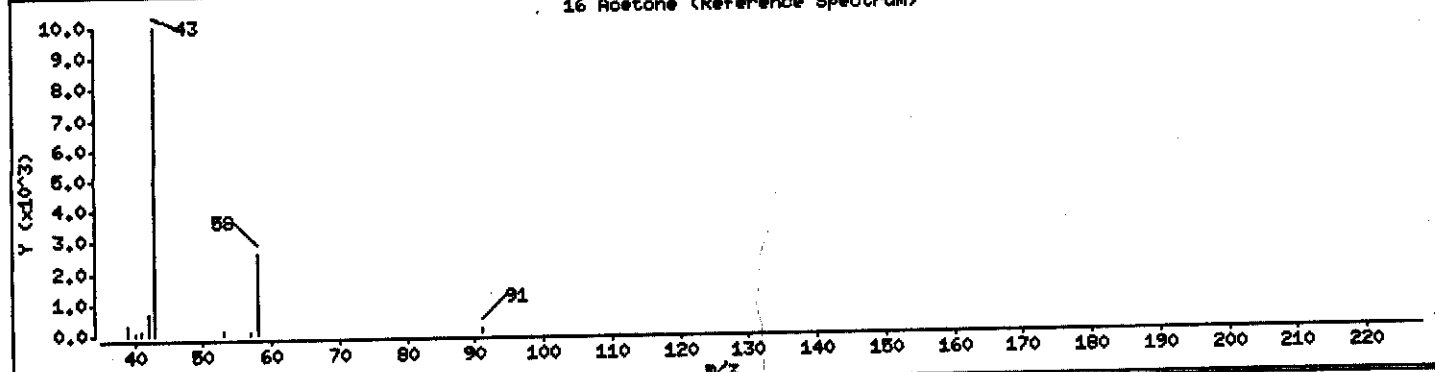
Scan 116 (2.746 min) of UXJ24288.D



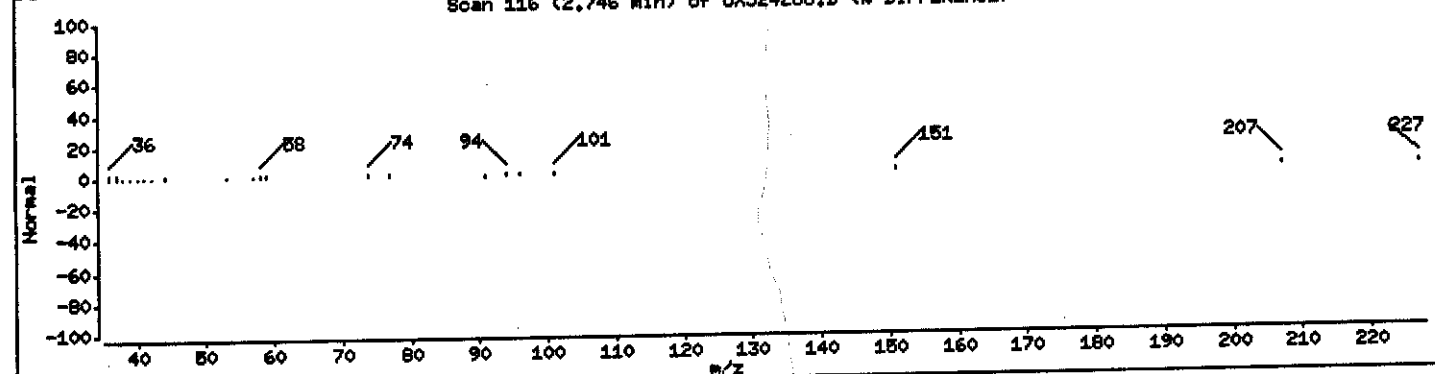
Scan 116 (2.746 min) of UXJ24288.D (Subtracted)



16 Acetone (Reference Spectrum)



Scan 116 (2.746 min) of UXJ24288.D (% DIFFERENCE)



Data File: \\qpanoh04\dd\chem\MSV\z3ux11.1\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:26

Client ID: VEB41/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: GRDX61AA,5ML/5ML

Purge Volume: 5.0

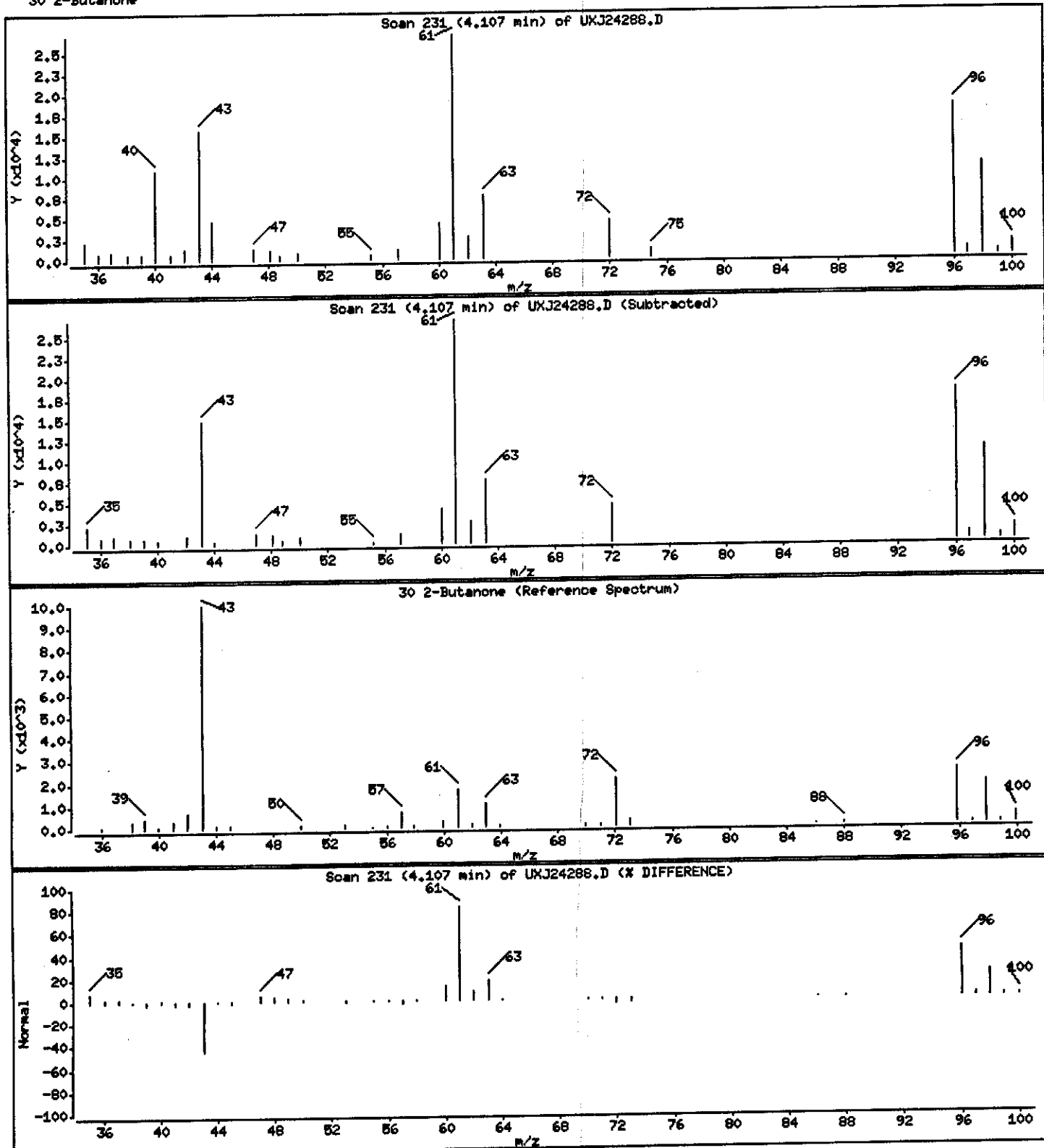
Operator: 43582

Column phase: DB624

Column diameter: 0.18

30 2-Butanone

Concentration: 2.160 ug/L



Data File: \\qcanno04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VEB41/26.5-31.5/092

Instrument: s3ux11.i

Sample Info: GRDX61AA,5ML/5ML

Purge Volume: 5.0

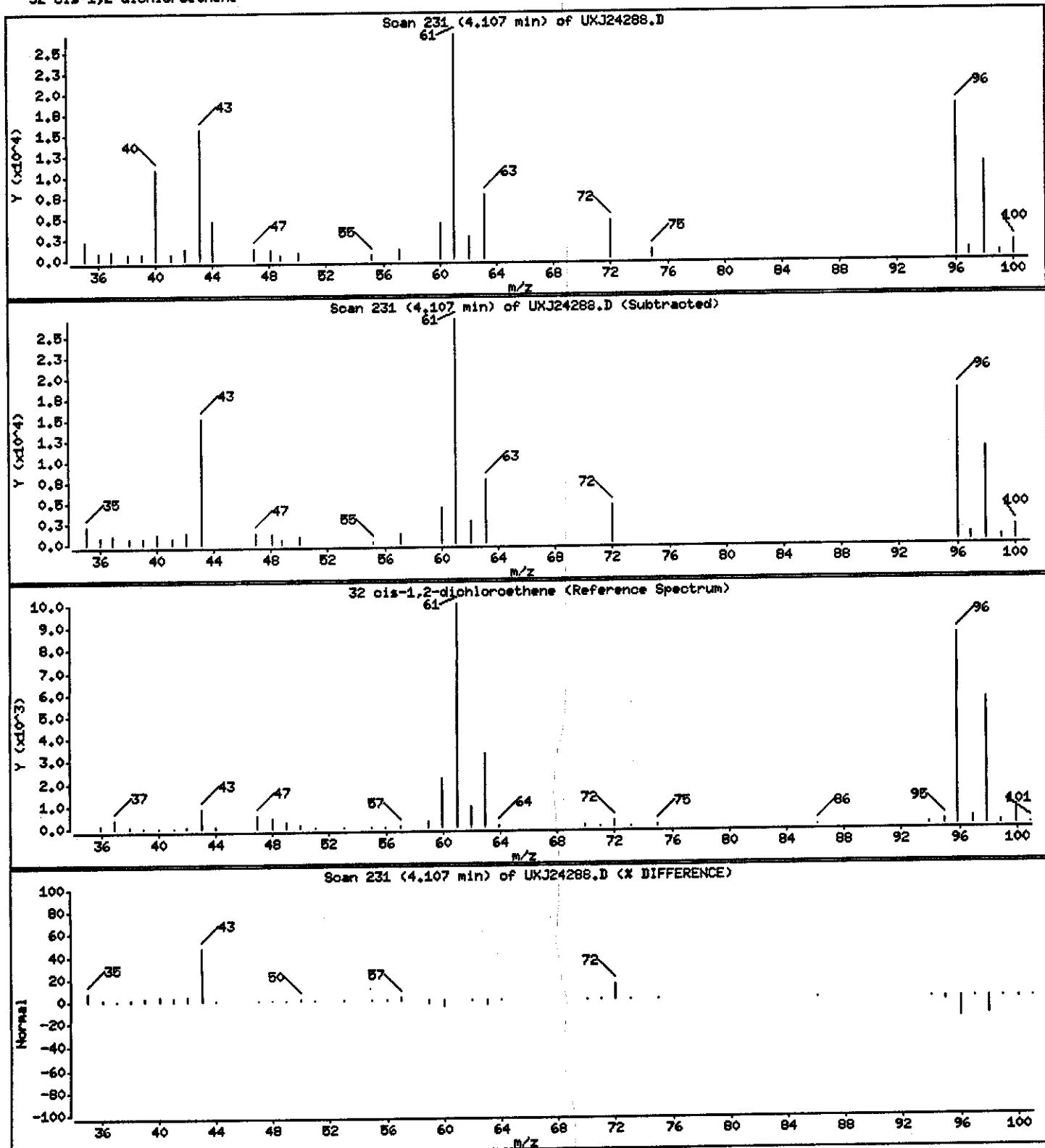
Operator: 43582

Column phase: DB624

Column diameter: 0.18

32 cis-1,2-dichloroethene

Concentration: 0.8998 ug/L



Data File: \\qoanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VEB41/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: CRDX61AA,5ML/5ML

Purge Volume: 5.0

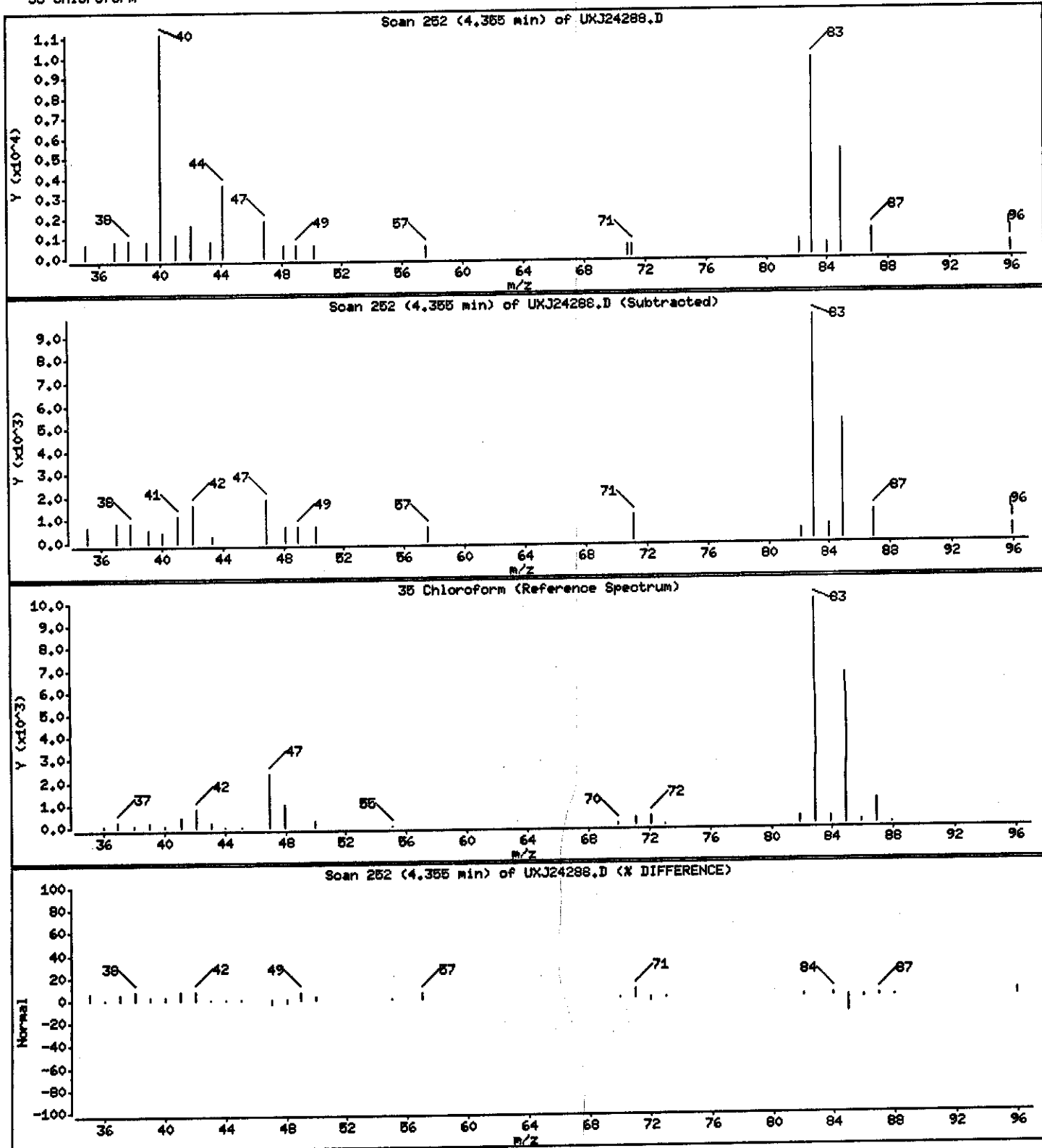
Operator: 43582

Column phase: DB624

Column diameter: 0.18

35 Chloroform

Concentration: 0.2654 ug/L



Data File: \\qcanoh04\dd\chem\MSI\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VE541/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: CRDX61AA,BML/BML

Purge Volume: 5.0

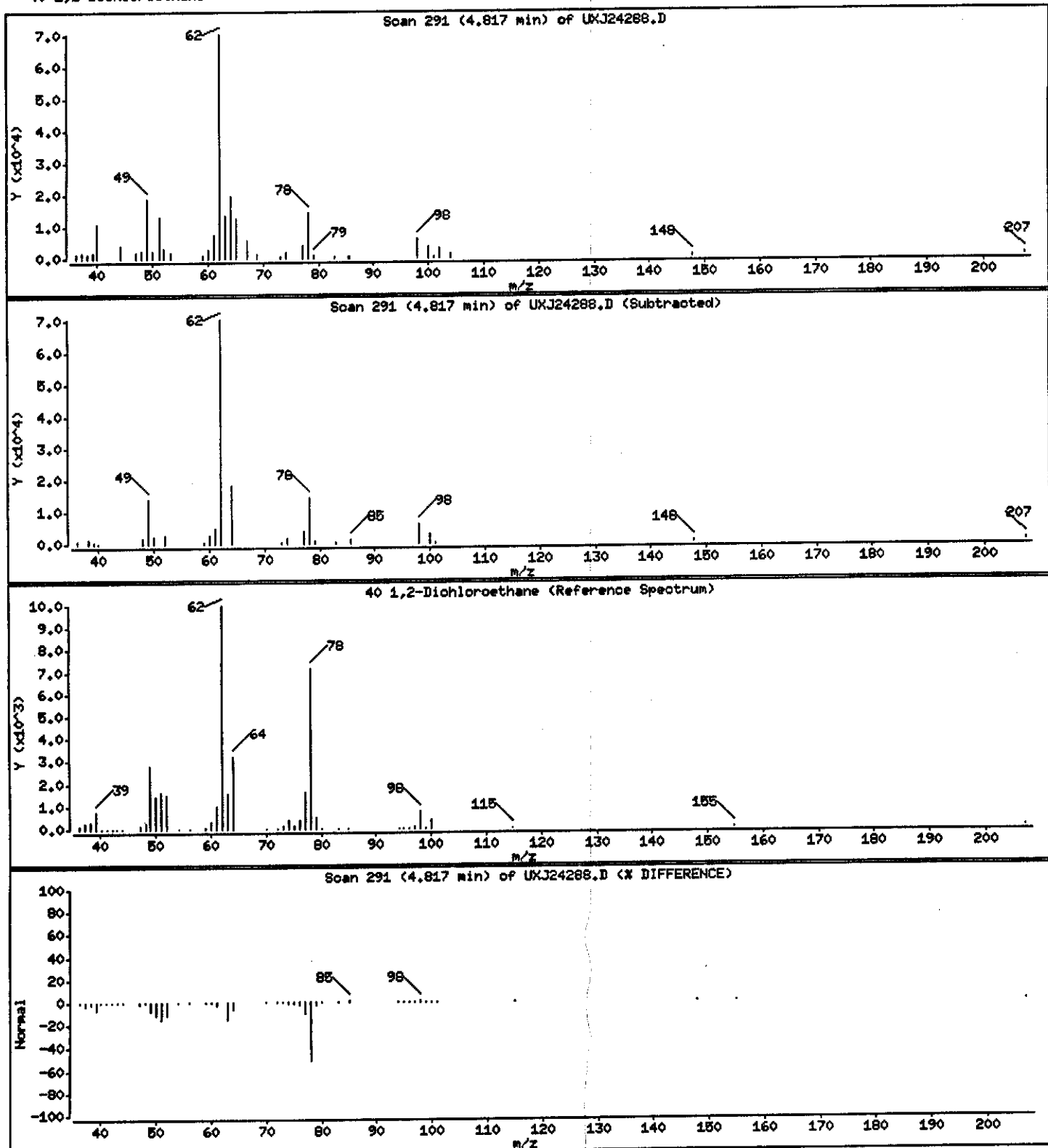
Operator: 43582

Column phase: DB624

Column diameter: 0.18

40 1,2-Dichloroethane

Concentration: 2.380 ug/L



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J41001A.b\\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VE541/26.5-31.5/092

Sample Info: GRDX61AA,5ML/5ML

Purge Volume: 5.0

Column phase: DB624

Instrument: a3ux11.i

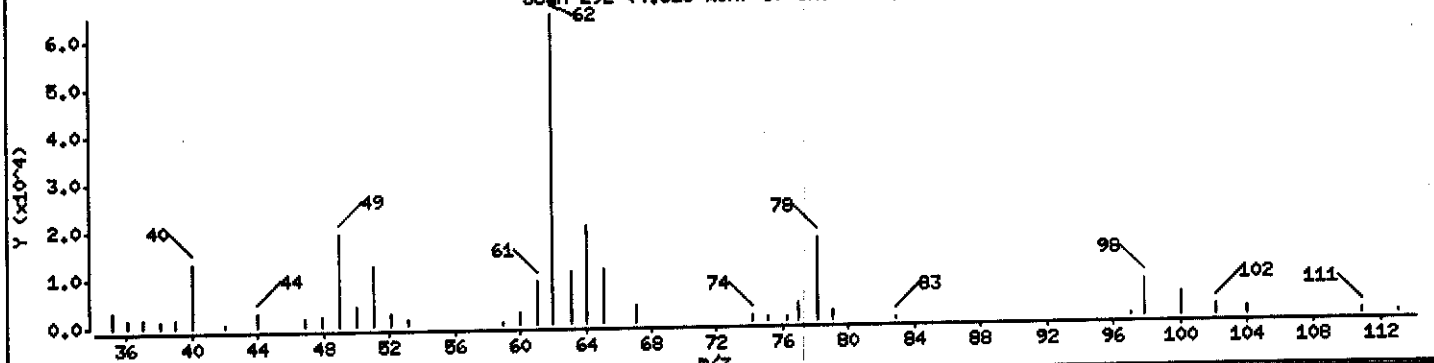
Operator: 43582

Column diameter: 0.18

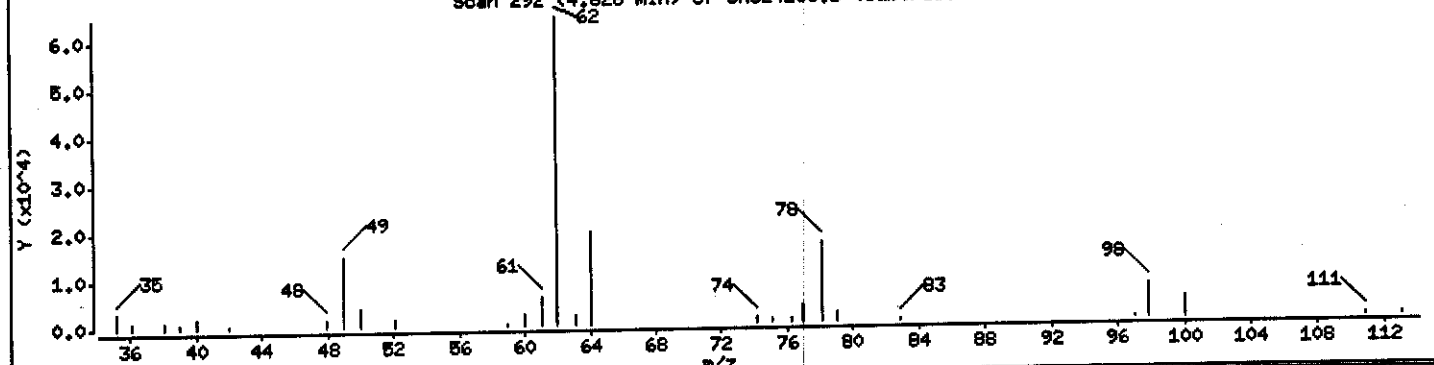
Concentration: 0.2367 ug/L

41 Benzene

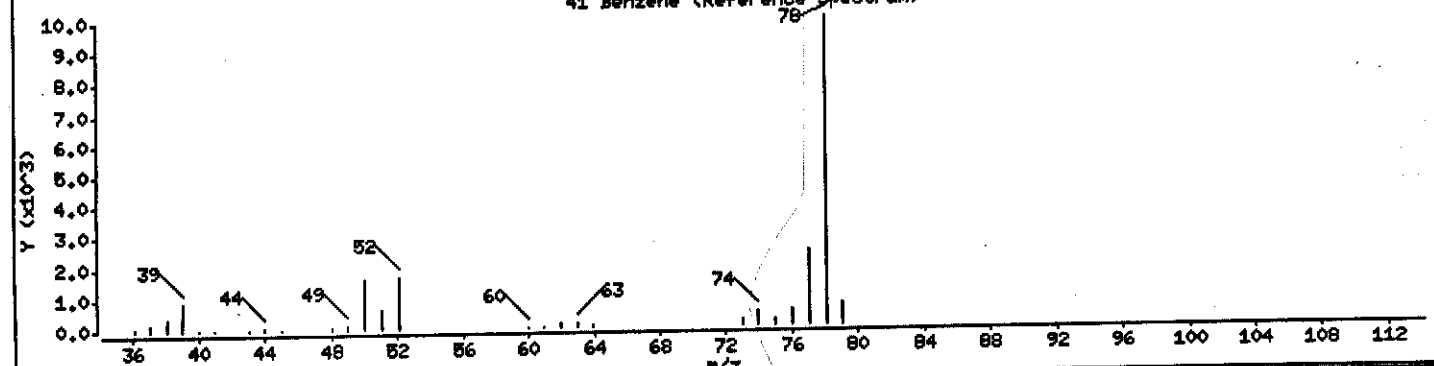
Scan 292 (4.828 min) of UXJ24288.D



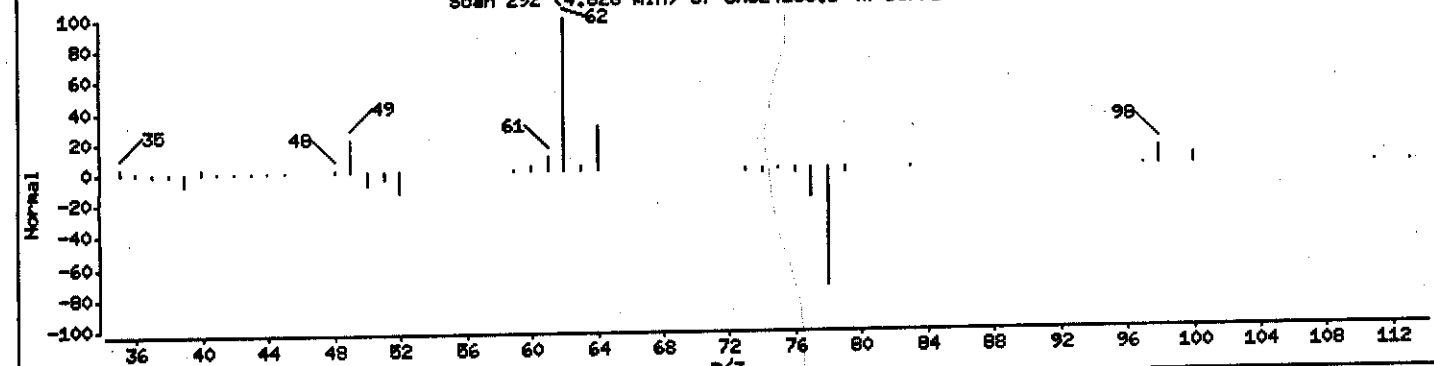
Scan 292 (4.828 min) of UXJ24288.D (Subtracted)



41 Benzene (Reference Spectrum)



Scan 292 (4.828 min) of UXJ24288.D (% DIFFERENCE)



Data File: \\qcarch04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VE541/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: GRDX61AA,5ML/5ML

Purge Volume: 5.0

Operator: 43582

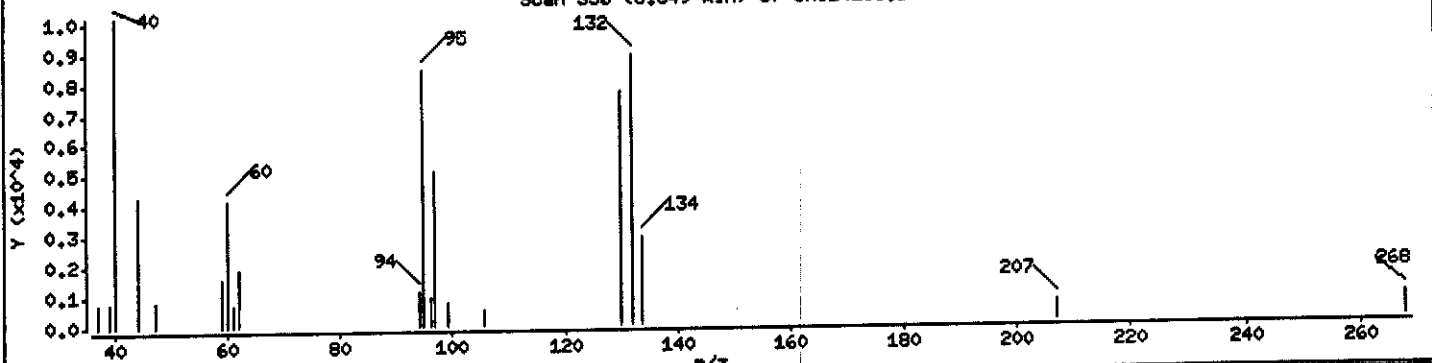
Column phase: DB624

Column diameter: 0.18

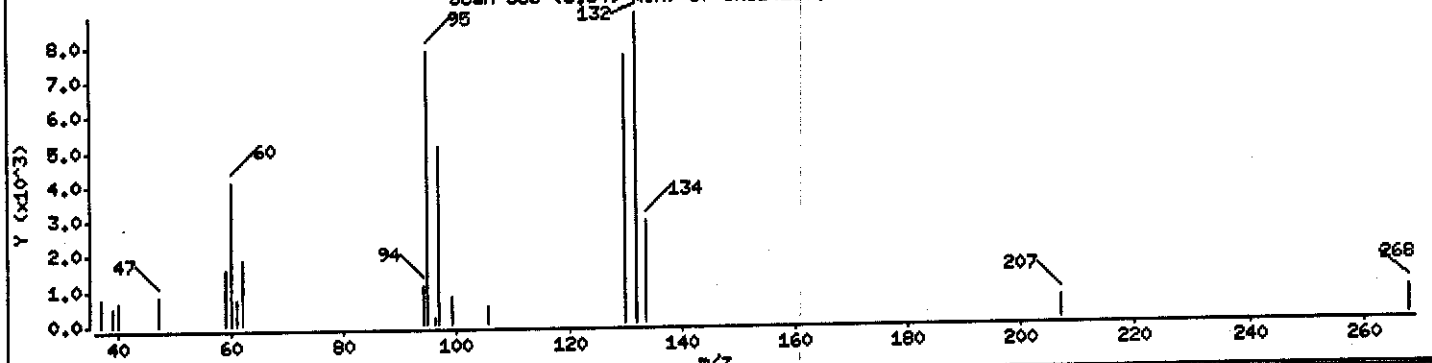
42 Trichloroethene

Concentration: 0.3780 ug/L

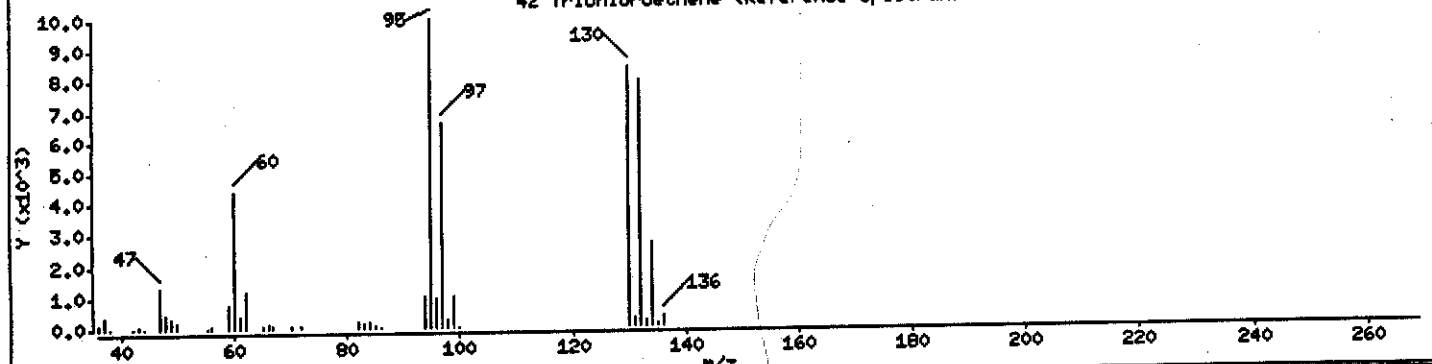
Scan 336 (5.349 min) of UXJ24288.D



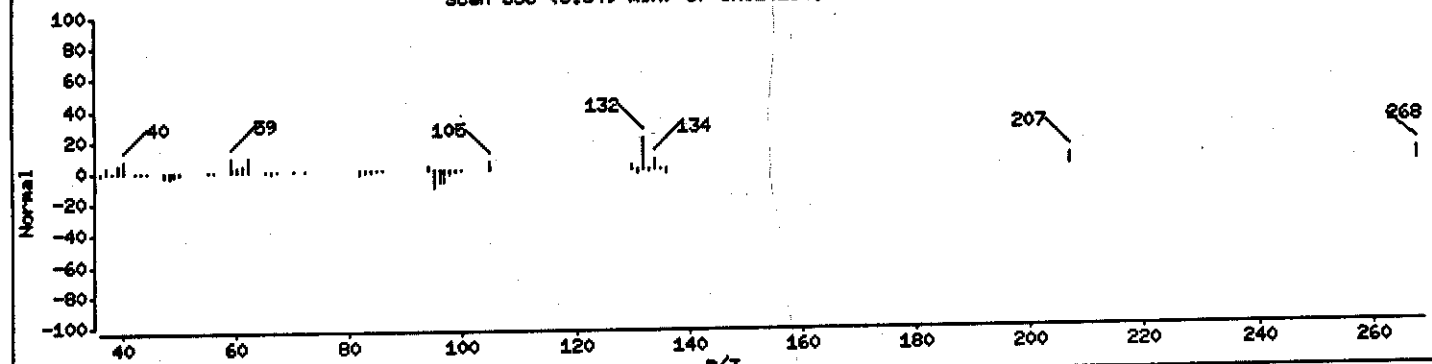
Scan 336 (5.349 min) of UXJ24288.D (Subtracted)



42 Trichloroethene (Reference Spectrum)



Scan 336 (5.349 min) of UXJ24288.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VE541/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: GRDX61AA,5ML/5ML

Operator: 43592

Purge Volume: 5.0

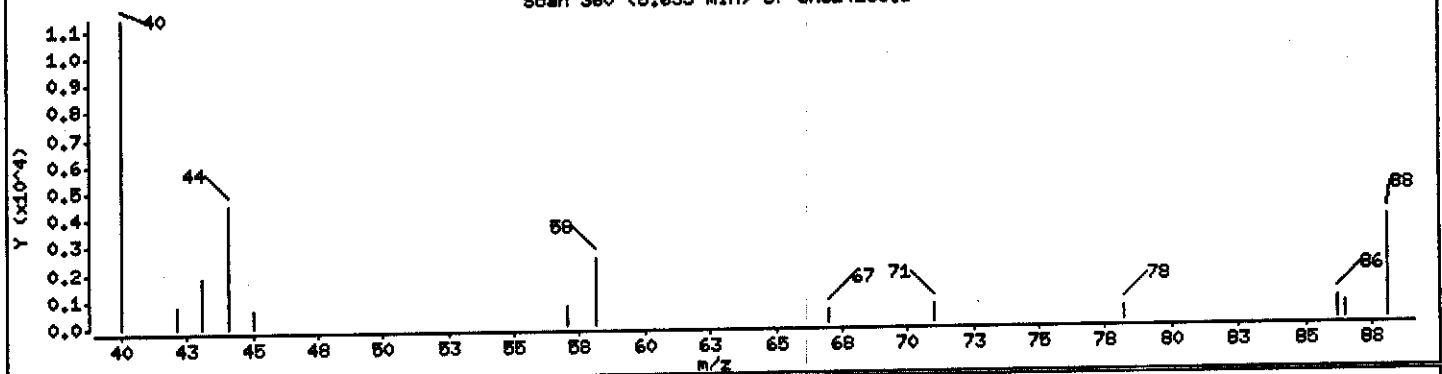
Column diameter: 0.18

Column phase: DB624

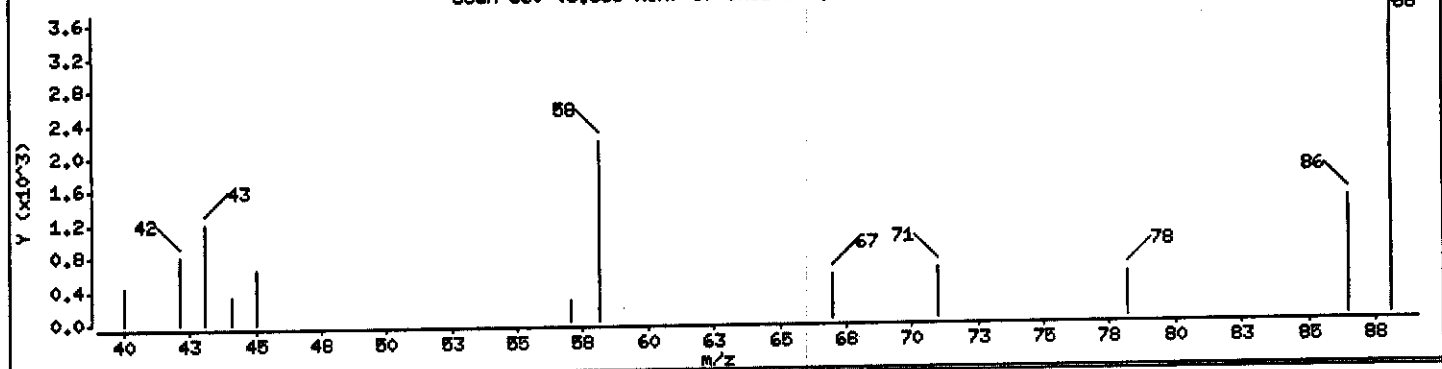
Concentration: 19.265 ug/L

44 1,4-Dioxane

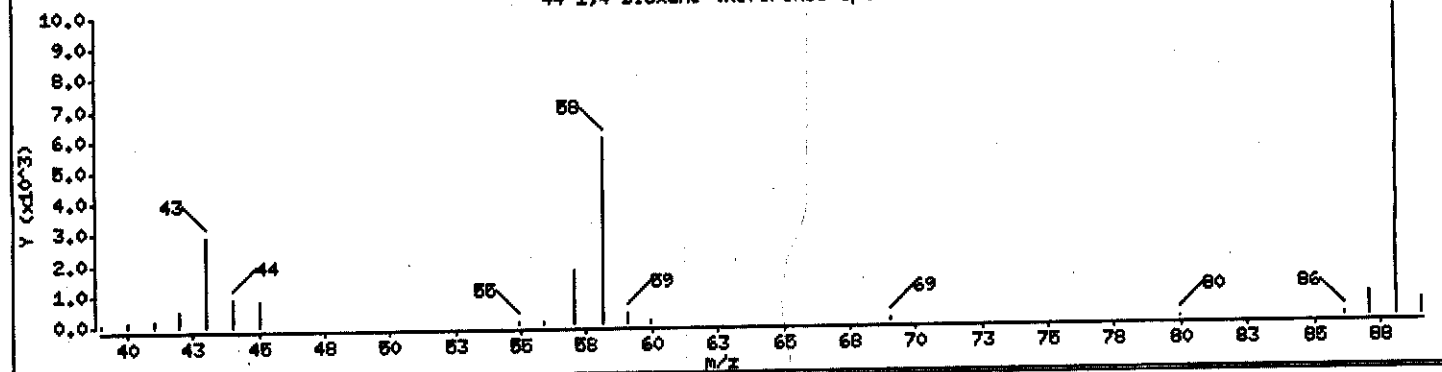
Scan 360 (5.633 min) of UXJ24288.D



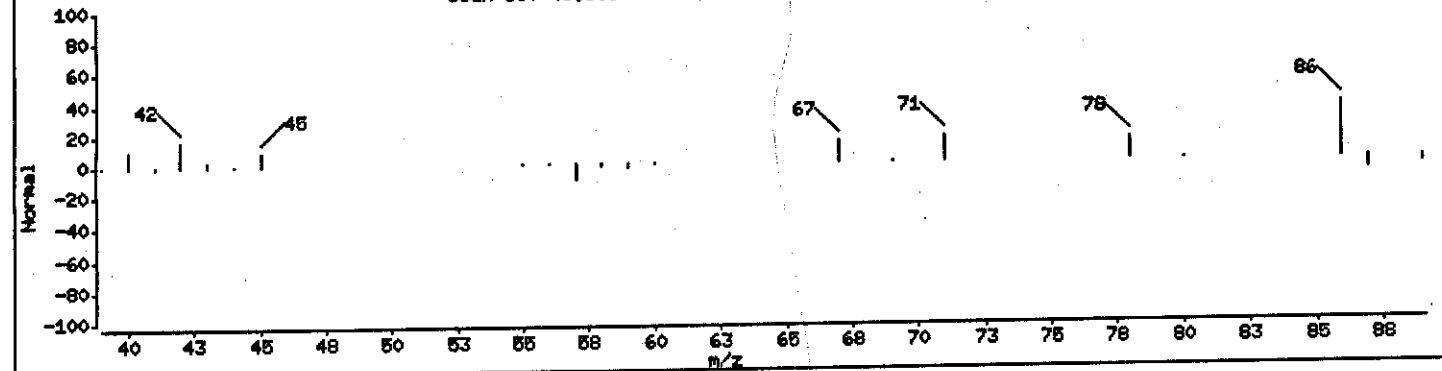
Scan 360 (5.633 min) of UXJ24288.D (Subtracted)



44 1,4-Dioxane (Reference Spectrum)



Scan 360 (5.633 min) of UXJ24288.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24288.D

Date : 01-OCT-2004 12:28

Client ID: VE841/26.5-31.5/092

Instrument: z3ux11.i

Sample Info: CRDX61AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

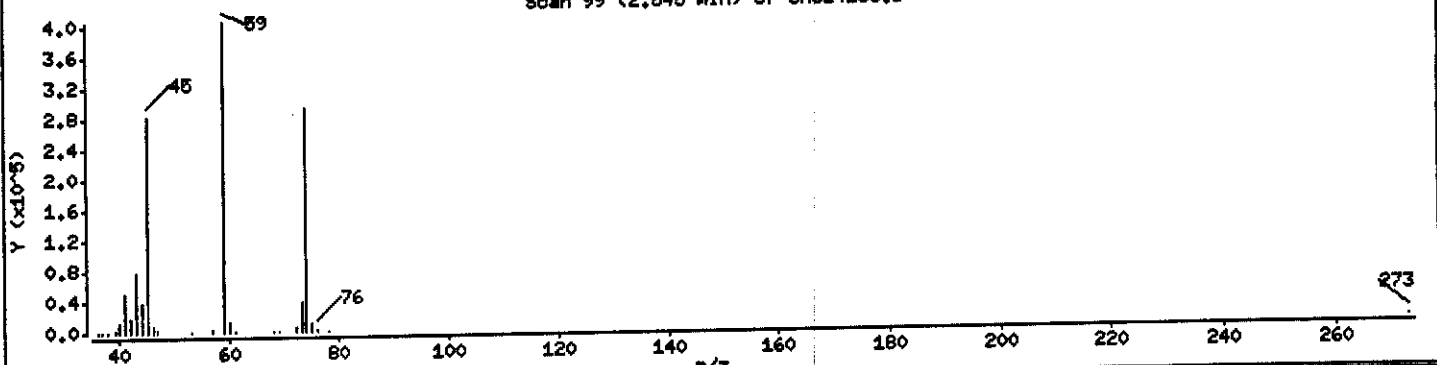
Column diameter: 0.18

Column phase: DB624

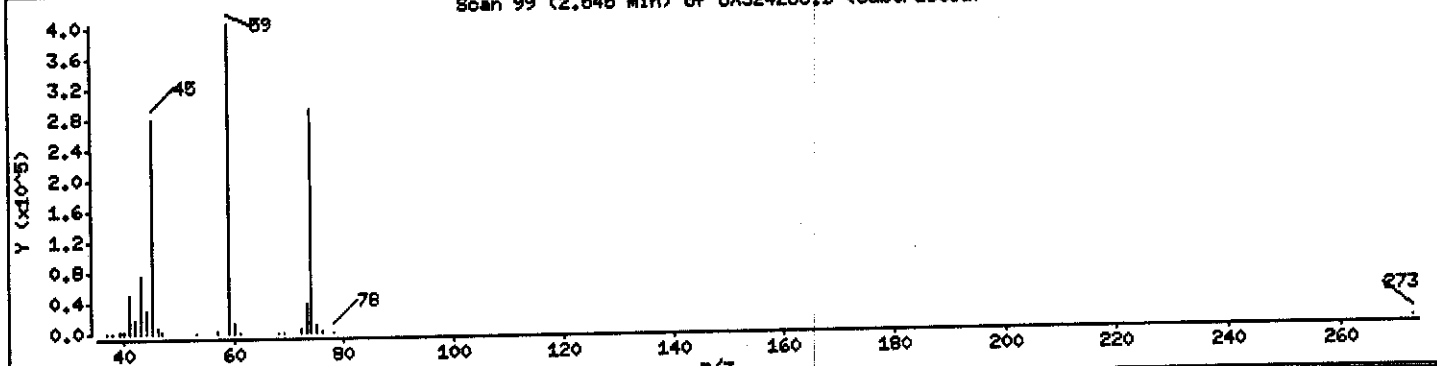
Concentration: 25.830 ug/L

89 Ethyl Ether

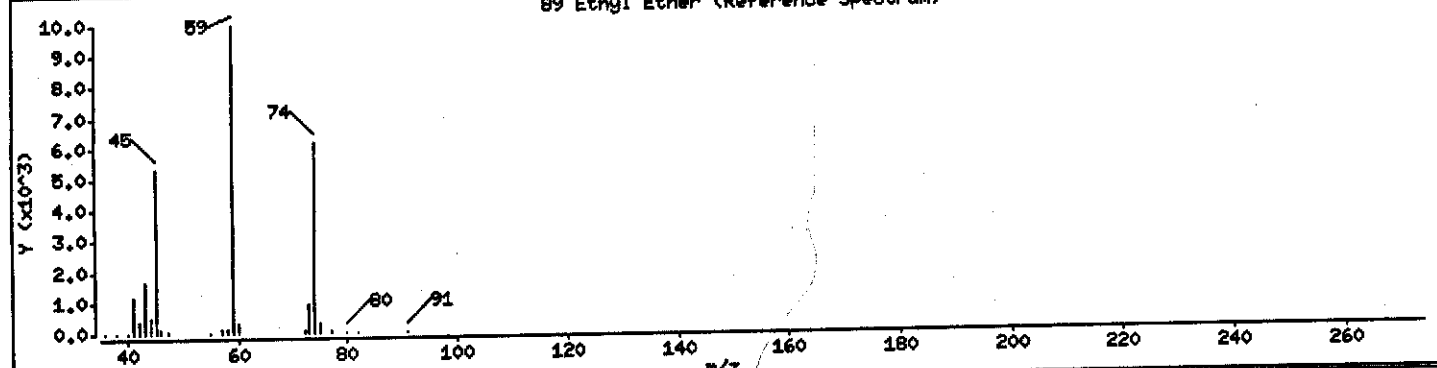
Scan 99 (2.545 min) of UXJ24288.D



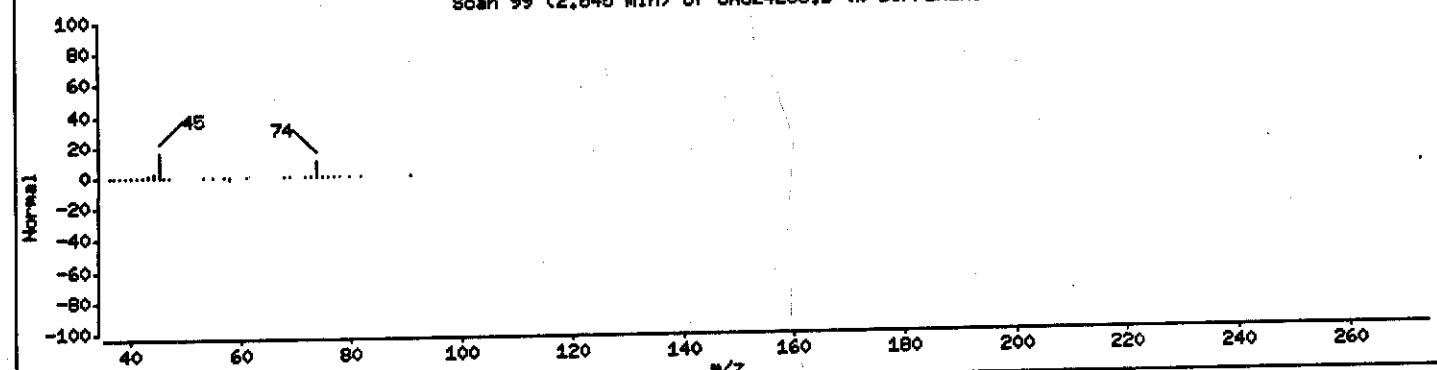
Scan 99 (2.545 min) of UXJ24288.D (Subtracted)



89 Ethyl Ether (Reference Spectrum)



Scan 99 (2.545 min) of UXJ24288.D (% DIFFERENCE)



PAYNE FIRM INC.

Client Sample ID: TRIP BLANK/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-004 Work Order #....: GRD0D1AA Matrix.....: WQ
 Date Sampled....: 09/28/04 Date Received...: 09/29/04
 Prep Date.....: 10/01/04 Analysis Date...: 10/01/04
 Prep Batch #....: 4275213
 Dilution Factor: 1 Initial Wgt/Vol.: 5 mL Final Wgt/Vol.: 5 mL
 Method.....: SW846 8260B

| PARAMETER | RESULT | REPORTING | |
|------------------------------------|----------|-----------|-------|
| | | LIMIT | UNITS |
| Acetone | 0.95 J | 10 | ug/L |
| Acetonitrile | ND | 20 | ug/L |
| Acrolein | ND | 20 | ug/L |
| Acrylonitrile | ND | 20 | ug/L |
| Benzene | ND | 1.0 | ug/L |
| Bromodichloromethane | ND | 1.0 | ug/L |
| Bromoform | ND | 1.0 | ug/L |
| Bromomethane | ND | 1.0 | ug/L |
| 2-Butanone | 0.48 J | 10 | ug/L |
| Carbon disulfide | ND | 1.0 | ug/L |
| Carbon tetrachloride | ND | 1.0 | ug/L |
| Chlorobenzene | ND | 1.0 | ug/L |
| Chloroprene | ND | 2.0 | ug/L |
| Dibromochloromethane | ND | 1.0 | ug/L |
| Chloroethane | ND | 1.0 | ug/L |
| Chloroform | ND | 1.0 | ug/L |
| Chloromethane | 0.22 J,B | 1.0 | ug/L |
| 3-Chloropropene | ND | 2.0 | ug/L |
| 1,2-Dibromo-3-chloropropane (DBCP) | ND | 2.0 | ug/L |
| 1,2-Dibromoethane | ND | 1.0 | ug/L |
| Dibromomethane | ND | 1.0 | ug/L |
| trans-1,4-Dichloro-2-butene | ND | 1.0 | ug/L |
| 1,1-Dichloroethane | ND | 1.0 | ug/L |
| 1,2-Dichloroethane | ND | 1.0 | ug/L |
| cis-1,2-Dichloroethene | ND | 1.0 | ug/L |
| trans-1,2-Dichloroethene | ND | 1.0 | ug/L |
| 1,1-Dichloroethene | ND | 1.0 | ug/L |
| 1,2-Dichloroethene (total) | ND | 2.0 | ug/L |
| Dichlorofluoromethane | ND | 2.0 | ug/L |
| 1,2-Dichloropropane | ND | 1.0 | ug/L |
| cis-1,3-Dichloropropene | ND | 1.0 | ug/L |
| trans-1,3-Dichloropropene | ND | 1.0 | ug/L |
| 1,4-Dioxane | ND | 50 | ug/L |
| Ethylbenzene | ND | 1.0 | ug/L |
| Ethyl methacrylate | ND | 1.0 | ug/L |

(Continued on next page)

PAYNE FIRM INC.

Client Sample ID: TRIP BLANK/092804

GC/MS Volatiles

Lot-Sample #....: A4I290193-004 Work Order #....: GRD0D1AA Matrix.....: WQ

| <u>PARAMETER</u> | <u>RESULT</u> | <u>REPORTING</u> | |
|---------------------------|---------------|------------------|--------------|
| | | <u>LIMIT</u> | <u>UNITS</u> |
| 2-Hexanone | ND | 10 | ug/L |
| Iodomethane | ND | 1.0 | ug/L |
| Isobutanol | ND | 50 | ug/L |
| Methacrylonitrile | ND | 2.0 | ug/L |
| Methylene chloride | ND | 1.0 | ug/L |
| Methyl methacrylate | ND | 2.0 | ug/L |
| 4-Methyl-2-pentanone | ND | 10 | ug/L |
| Propionitrile | ND | 4.0 | ug/L |
| Styrene | ND | 1.0 | ug/L |
| 1,1,1,2-Tetrachloroethane | ND | 1.0 | ug/L |
| 1,1,2,2-Tetrachloroethane | ND | 1.0 | ug/L |
| Tetrachloroethene | ND | 1.0 | ug/L |
| Toluene | 0.19 J | 1.0 | ug/L |
| 1,1,1-Trichloroethane | ND | 1.0 | ug/L |
| 1,1,2-Trichloroethane | ND | 1.0 | ug/L |
| Trichloroethene | ND | 1.0 | ug/L |
| Trichlorofluoromethane | ND | 1.0 | ug/L |
| 1,2,3-Trichloropropane | ND | 1.0 | ug/L |
| Vinyl acetate | ND | 2.0 | ug/L |
| Vinyl chloride | ND | 1.0 | ug/L |
| Xylenes (total) | ND | 2.0 | ug/L |

| <u>SURROGATE</u> | <u>PERCENT</u> <u>RECOVERY</u> | <u>RECOVERY</u> | |
|-----------------------|-----------------------------------|-----------------|--|
| | | <u>LIMITS</u> | |
| Dibromofluoromethane | 118 | (73 - 122) | |
| 1,2-Dichloroethane-d4 | 111 | (61 - 128) | |
| Toluene-d8 | 87 | (76 - 110) | |
| 4-Bromofluorobenzene | 80 | (74 - 116) | |

NOTE(S) :

J Estimated result. Result is less than RL.

B Method blank contamination. The associated method blank contains the target analyte at a reportable level.

Data File: \\pcando4\dd\chen\HSV\aa3\ad1.i\\410019.b\\J24289.D

Date : 01-OCT-2004 12:51

Client ID: TRIP BLANK\092804

Sample Info: GRD0019A, EML/5FL

Purge Volume: 5.0

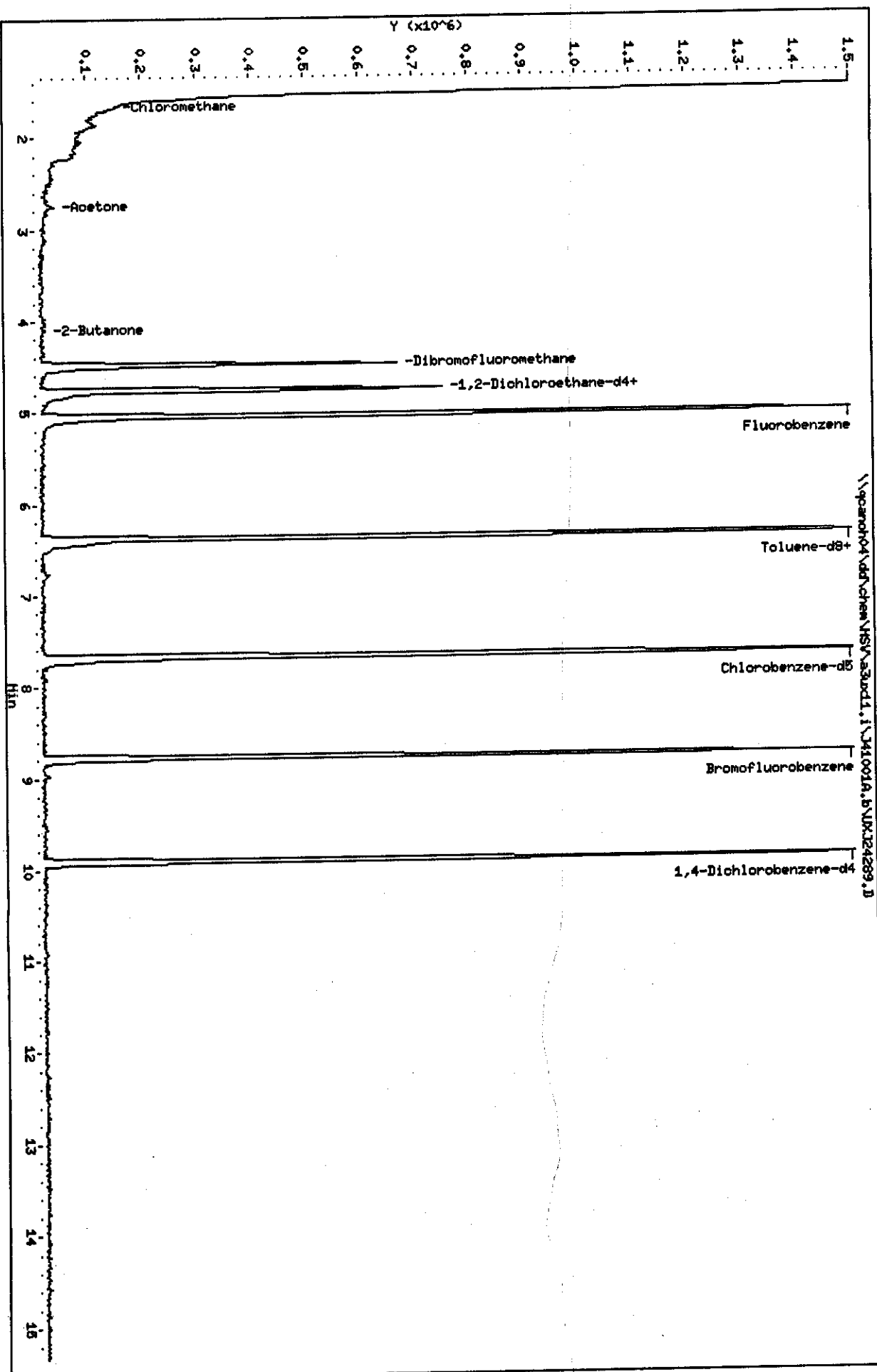
Column Phase: DB624

Instrument: a3ad1.i

Operator: 43882

Column diameter: 0.19

\\pcando4\dd\chen\HSV\aa3\ad1.i\\410019.b\\J24289.D



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24289.D
Report Date: 04-Oct-2004 10:02

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24289.D
Lab Smp Id: GRD0D1AA Client Smp ID: TRIP BLANK/092804
Inj Date : 01-OCT-2004 12:51
Operator : 43582 Inst ID: a3ux11.i
Smp Info : GRD0D1AA,5ML/5ML
Misc Info : J41001A,8260LLUX11,,43582
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 13
Dil Factor: 1.00000
Integrator: HP RTE
Target Version: 4.04
Processing Host: CANPMSV07
Compound Sublist: 4-8260+IX.sub

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|-----------|--------------------------|----------------|-------|------------------------|---------|----------|--------|---------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) | (ug/L) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 1774228 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1579425 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 650817 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 489194 | 59.2201 | 11.844 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 632748 | 55.7047 | 11.141 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 1645320 | 43.3617 | 8.672 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 644939 | 40.0034 | 8.001 | |
| 8 | Dichlorodifluoromethane | 85 | | Compound Not Detected. | | | | |
| 9 | Chloromethane | 50 | 1.669 | 1.704 (0.331) | 18194 | 1.09858 | 0.2197 | |
| 10 | Vinyl Chloride | 62 | | Compound Not Detected. | | | | |
| 11 | Bromomethane | 94 | | Compound Not Detected. | | | | |
| 12 | Chloroethane | 64 | | Compound Not Detected. | | | | |
| 13 | Trichlorofluoromethane | 101 | | Compound Not Detected. | | | | |
| 15 | Acrolein | 56 | | Compound Not Detected. | | | | |
| 16 | Acetone | 43 | 2.745 | 2.745 (0.545) | 38304 | 4.76023 | 0.9520 | |
| 17 | 1,1-Dichloroethene | 96 | | Compound Not Detected. | | | | |
| 18 | Freon-113 | 151 | | Compound Not Detected. | | | | |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|---------------------------------|-----------|------|----------------|---------------|--------|------------------------|-------------------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 19 Iodomethane | | 142 | | | | Compound Not Detected. | |
| 20 Carbon Disulfide | | 76 | | | | Compound Not Detected. | |
| 21 Methylene Chloride | | 84 | | | | Compound Not Detected. | |
| 22 Acetonitrile | | 41 | | | | Compound Not Detected. | |
| 23 Acrylonitrile | | 53 | | | | Compound Not Detected. | |
| 24 Methyl tert-butyl ether | | 73 | | | | Compound Not Detected. | |
| 25 trans-1,2-Dichloroethene | | 96 | | | | Compound Not Detected. | |
| 26 Hexane | | 86 | | | | Compound Not Detected. | |
| 27 Vinyl acetate | | 43 | | | | Compound Not Detected. | |
| 28 1,1-Dichloroethane | | 63 | | | | Compound Not Detected. | |
| 29 tert-Butyl Alcohol | | 59 | | | | Compound Not Detected. | |
| 30 2-Butanone | | 43 | 4.094 | 4.106 (0.812) | 11549 | 2.37653 | 0.4753 |
| M 31 1,2-Dichloroethene (total) | | 96 | | | | Compound Not Detected. | |
| 32 cis-1,2-dichloroethene | | 96 | | | | Compound Not Detected. | |
| 33 2,2-Dichloropropane | | 77 | | | | Compound Not Detected. | |
| 34 Bromochloromethane | | 128 | | | | Compound Not Detected. | |
| 35 Chloroform | | 83 | | | | Compound Not Detected. | |
| 36 Tetrahydrofuran | | 42 | | | | Compound Not Detected. | |
| 37 1,1,1-Trichloroethane | | 97 | | | | Compound Not Detected. | |
| 38 1,1-Dichloropropene | | 75 | | | | Compound Not Detected. | |
| 39 Carbon Tetrachloride | | 117 | | | | Compound Not Detected. | |
| 40 1,2-Dichloroethane | | 62 | | | | Compound Not Detected. | |
| 41 Benzene | | 78 | 4.828 | 4.828 (0.958) | 37585 | 0.92011 | 0.1840 |
| 42 Trichloroethene | | 130 | | | | Compound Not Detected. | |
| 43 1,2-Dichloropropane | | 63 | | | | Compound Not Detected. | |
| 44 1,4-Dioxane | | 88 | | | | Compound Not Detected. | |
| 45 Dibromomethane | | 93 | | | | Compound Not Detected. | |
| 46 Bromodichloromethane | | 83 | | | | Compound Not Detected. | |
| 47 2-Chloroethyl vinyl ether | | 63 | | | | Compound Not Detected. | |
| 48 cis-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | |
| 49 4-Methyl-2-pentanone | | 43 | | | | Compound Not Detected. | |
| 50 Toluene | | 91 | 6.437 | 6.437 (0.838) | 44013 | 0.93307 | 0.1866 |
| 51 trans-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | |
| 52 Ethyl Methacrylate | | 69 | | | | Compound Not Detected. | |
| 53 1,1,2-Trichloroethane | | 97 | | | | Compound Not Detected. | |
| 54 1,3-Dichloropropane | | 76 | | | | Compound Not Detected. | |
| 55 Tetrachloroethene | | 164 | | | | Compound Not Detected. | |
| 56 2-Hexanone | | 43 | | | | Compound Not Detected. | |
| 57 Dibromochloromethane | | 129 | | | | Compound Not Detected. | |
| 58 1,2-Dibromoethane | | 107 | | | | Compound Not Detected. | |
| 59 Chlorobenzene | | 112 | | | | Compound Not Detected. | |
| 60 1,1,1,2-Tetrachloroethane | | 131 | | | | Compound Not Detected. | |
| 61 Ethylbenzene | | 106 | | | | Compound Not Detected. | |
| 62 m + p-Xylene | | 106 | | | | Compound Not Detected. | |
| M 63 Xylenes (total) | | 106 | | | | Compound Not Detected. | |
| 64 Xylene-o | | 106 | | | | Compound Not Detected. | |
| 65 Styrene | | 104 | | | | Compound Not Detected. | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24289.D
Report Date: 04-Oct-2004 10:02

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | ON-COLUMN (ng) | FINAL (ug/L) |
|--------------------------------|-----------|------|----------------|--------|--------|------------------------|--------------------|------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | | |
| 66 Bromoform | ----- | 173 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 67 Isopropylbenzene | ----- | 105 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 68 1,1,2,2-Tetrachloroethane | ----- | 83 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 69 1,4-Dichloro-2-butene | ----- | 53 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 70 1,2,3-Trichloropropane | ----- | 110 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 71 Bromobenzene | ----- | 156 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 72 n-Propylbenzene | ----- | 120 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 73 2-Chlorotoluene | ----- | 126 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 74 1,3,5-Trimethylbenzene | ----- | 105 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 75 4-Chlorotoluene | ----- | 126 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 76 tert-Butylbenzene | ----- | 119 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 77 1,2,4-Trimethylbenzene | ----- | 105 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 78 sec-Butylbenzene | ----- | 105 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 79 4-Isopropyltoluene | ----- | 119 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 80 1,3-Dichlorobenzene | ----- | 146 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 81 1,4-Dichlorobenzene | ----- | 146 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 82 n-Butylbenzene | ----- | 91 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 83 1,2-Dichlorobenzene | ----- | 146 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 84 1,2-Dibromo-3-chloropropane | ----- | 157 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 85 1,2,4-Trichlorobenzene | ----- | 180 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 86 Hexachlorobutadiene | ----- | 225 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 87 Naphthalene | ----- | 128 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 88 1,2,3-Trichlorobenzene | ----- | 180 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 14 Dichlorofluoromethane | ----- | 67 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 89 Ethyl Ether | ----- | 59 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 91 3-Chloropropene | ----- | 76 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 92 Isopropyl Ether | ----- | 87 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 93 2-Chloro-1,3-butadiene | ----- | 53 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 94 Propionitrile | ----- | 54 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 95 Ethyl Acetate | ----- | 43 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 96 Methacrylonitrile | ----- | 41 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 97 Isobutanol | ----- | 41 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 99 n-Butanol | ----- | 56 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 100 Methyl Methacrylate | ----- | 41 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 101 2-Nitropropane | ----- | 41 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 103 Cyclohexanone | ----- | 55 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 98 Cyclohexane | ----- | 56 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 143 Methyl Acetate | ----- | 43 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 144 Methylcyclohexane | ----- | 83 | --- | --- | --- | Compound Not Detected. | ----- | ----- |
| 141 1,3,5-Trichlorobenzene | ----- | 180 | --- | --- | --- | Compound Not Detected. | ----- | ----- |

Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24289.D

Date : 01-OCT-2004 12:51

Client ID: TRIP BLANK/092804

Sample Info: GRDOD1AA,5ML/5ML

Purge Volume: 5.0

Column phase: DB624

Instrument: s3ux11.i

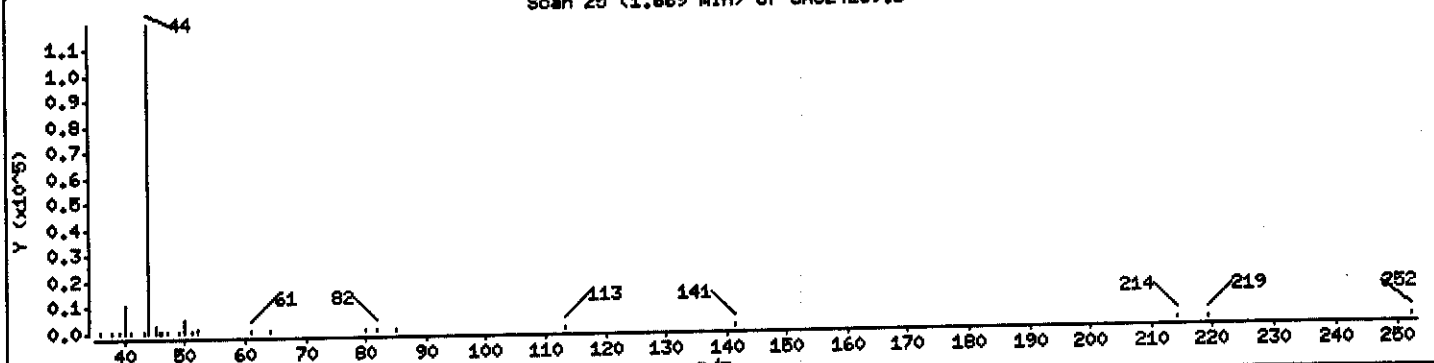
Operator: 43582

Column diameter: 0.18

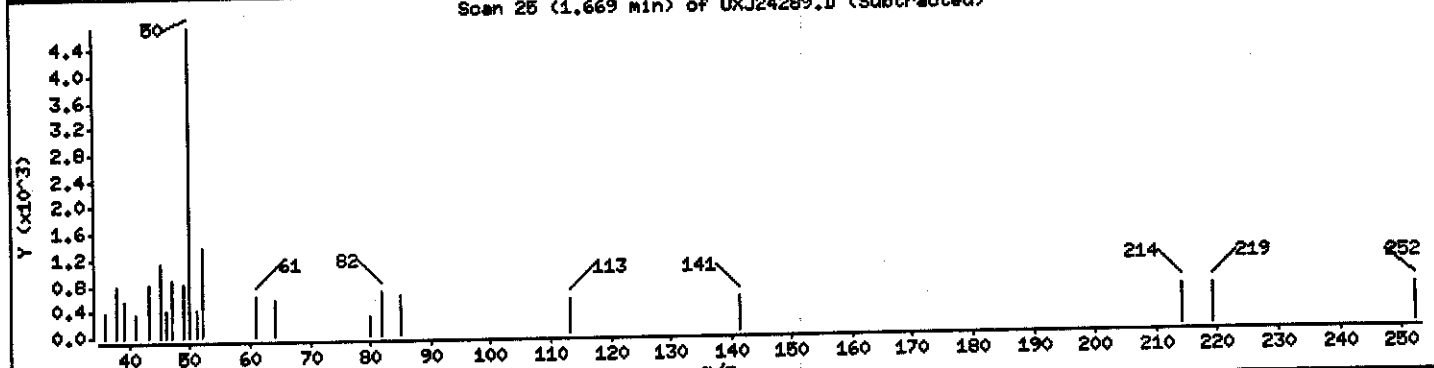
Concentration: 0.2197 ug/L

9 Chloromethane

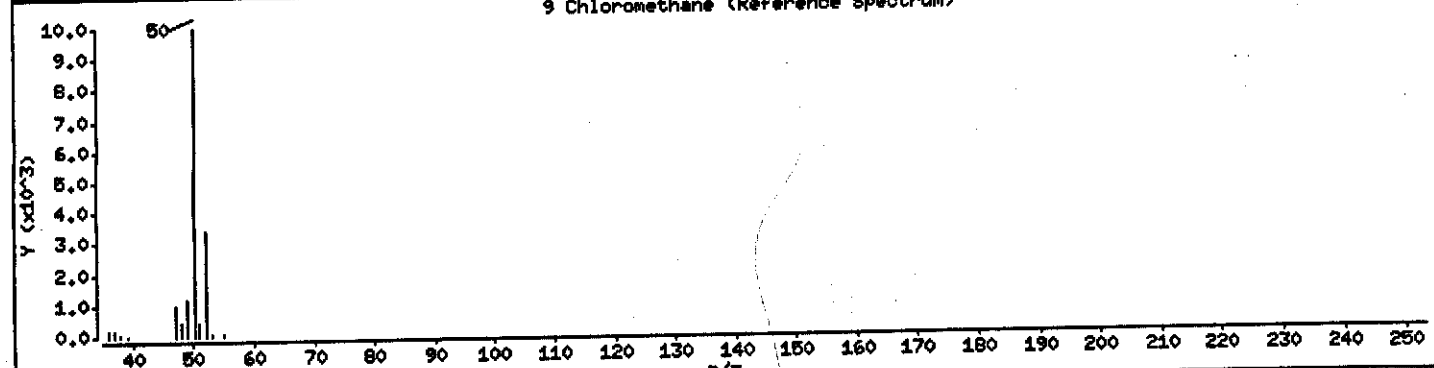
Scan 25 (1.669 min) of UXJ24289.D



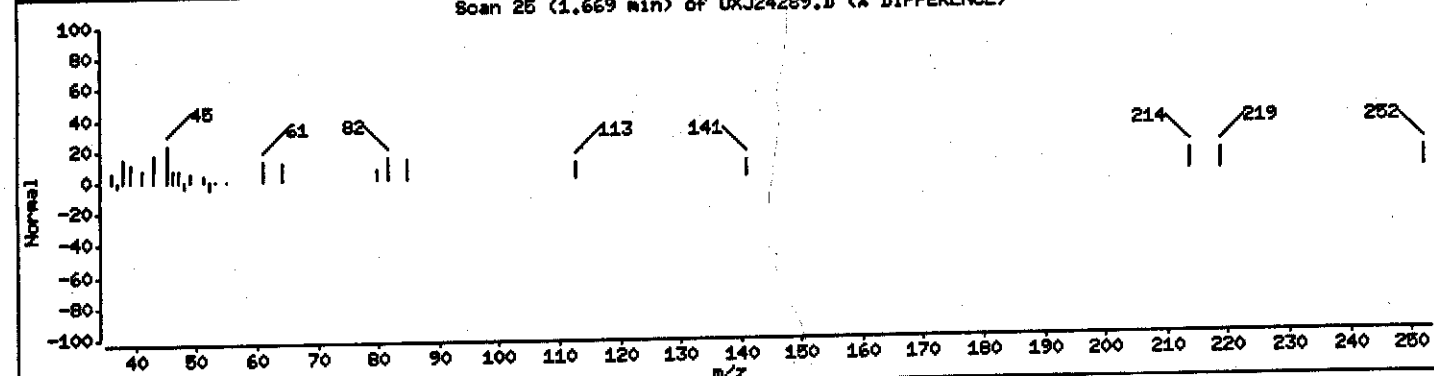
Scan 25 (1.669 min) of UXJ24289.D (Subtracted)



9 Chloromethane (Reference Spectrum)



Scan 25 (1.669 min) of UXJ24289.D (% DIFFERENCE)



Data File: \\qcanch04\dd\chem\MSV\z3ux11.i\J41001A.b\UXJ24289.D

Date : 01-OCT-2004 12:51

Client ID: TRIP BLANK/092804

Instrument: z3ux11.i

Sample Info: GRD001AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

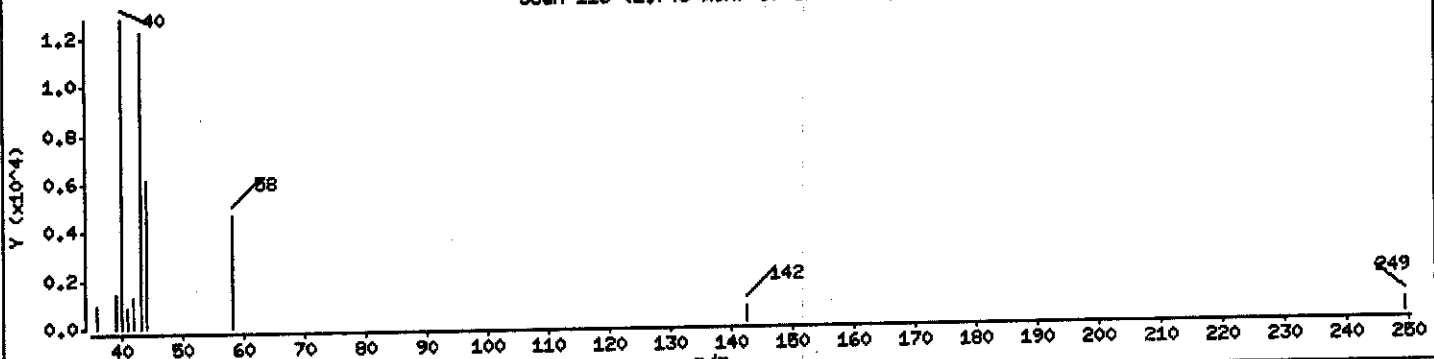
Column diameter: 0.18

Column phase: DB624

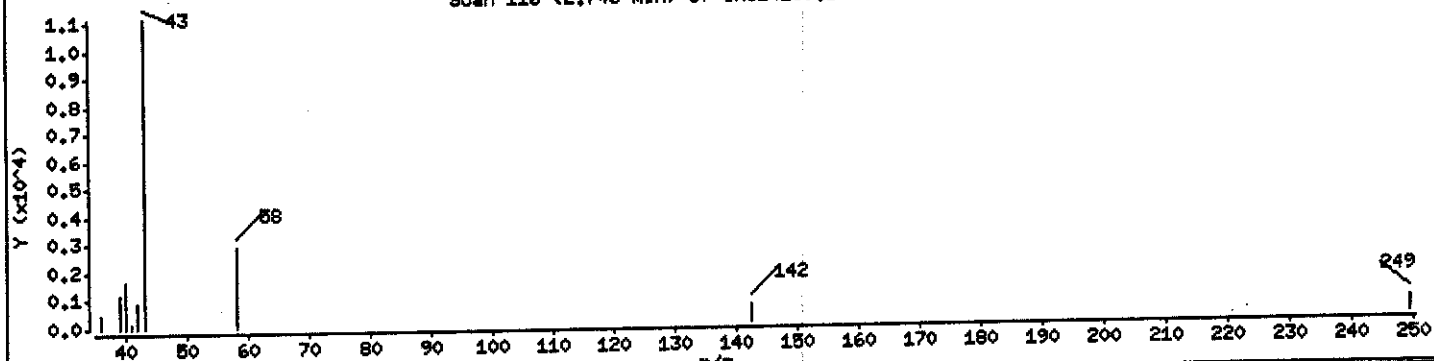
Concentration: 0.9520 ug/L

16 Acetone

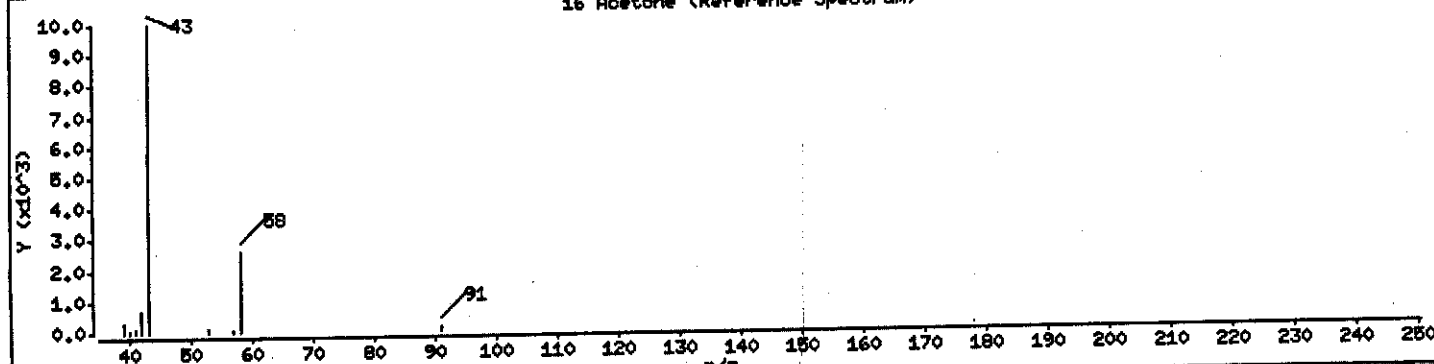
Scan 116 (2.746 min) of UXJ24289.D



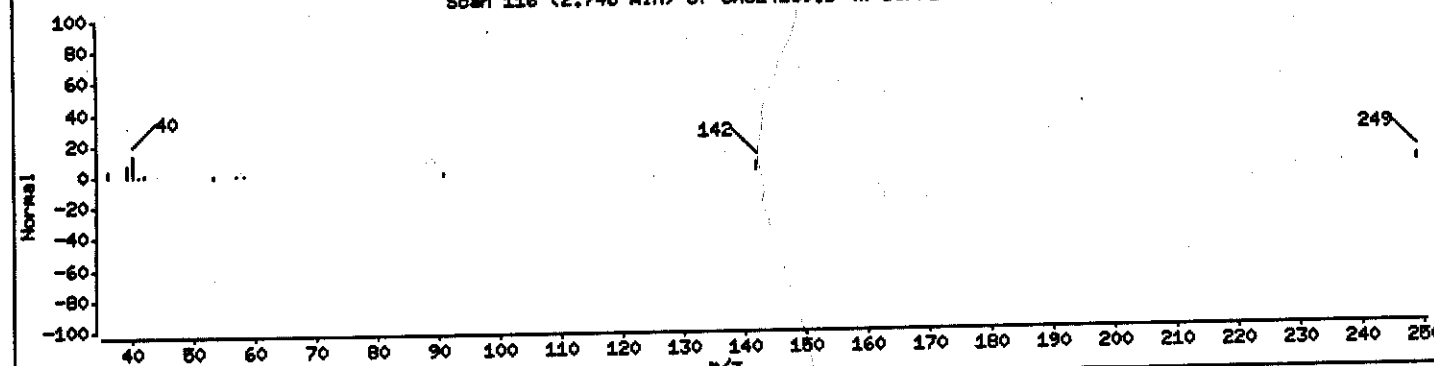
Scan 116 (2.746 min) of UXJ24289.D (Subtracted)



16 Acetone (Reference Spectrum)



Scan 116 (2.746 min) of UXJ24289.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.1\J41001A.b\UXJ24289.D

Date : 01-OCT-2004 12:51

Client ID: TRIP BLANK/092804

Sample Info: GRDOD1AA,5ML/5ML

Purge Volume: 5.0

Column phase: DB624

Instrument: z3ux11.1

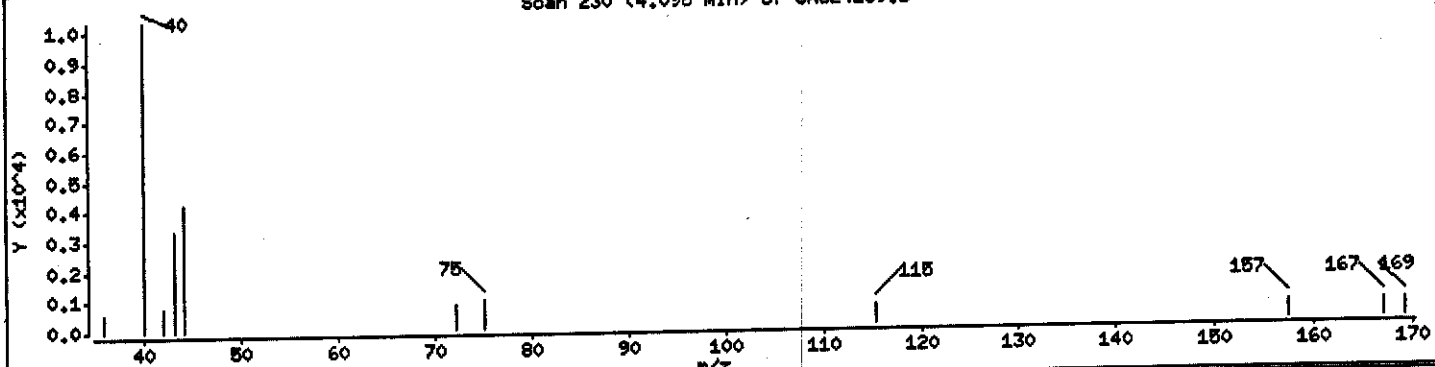
Operator: 43562

Column diameter: 0.18

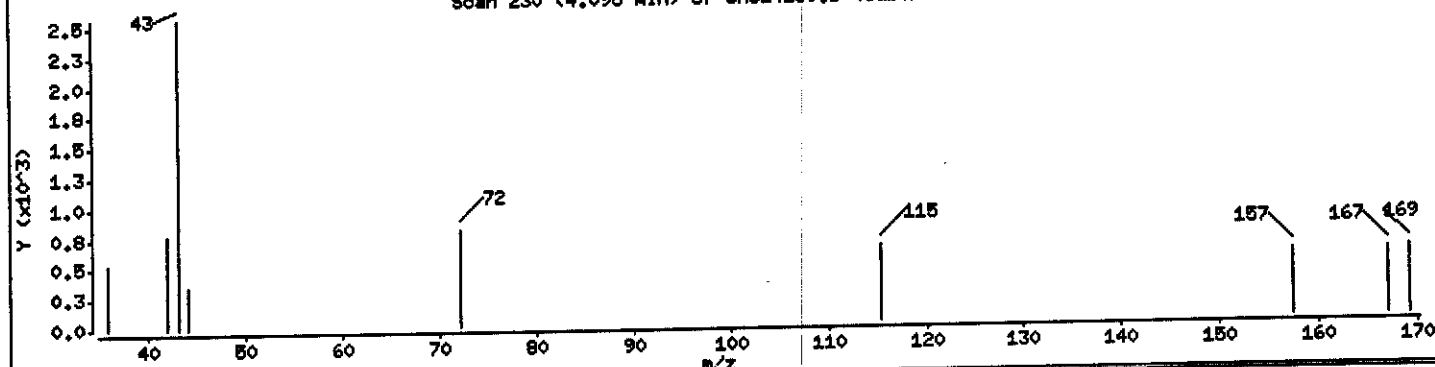
Concentration: 0,4753 ug/L

30 2-Butanone

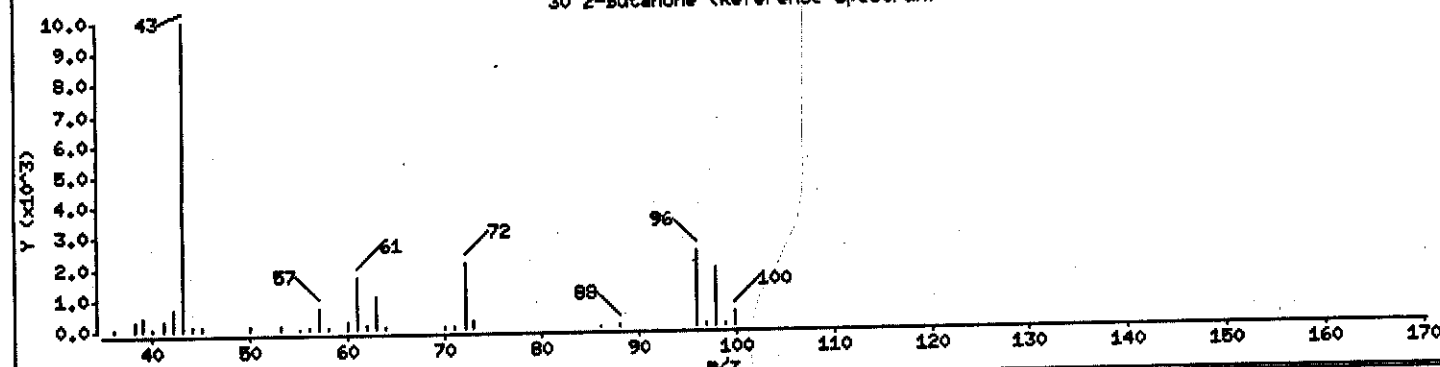
Scan 230 (4.095 min) of UXJ24289.D



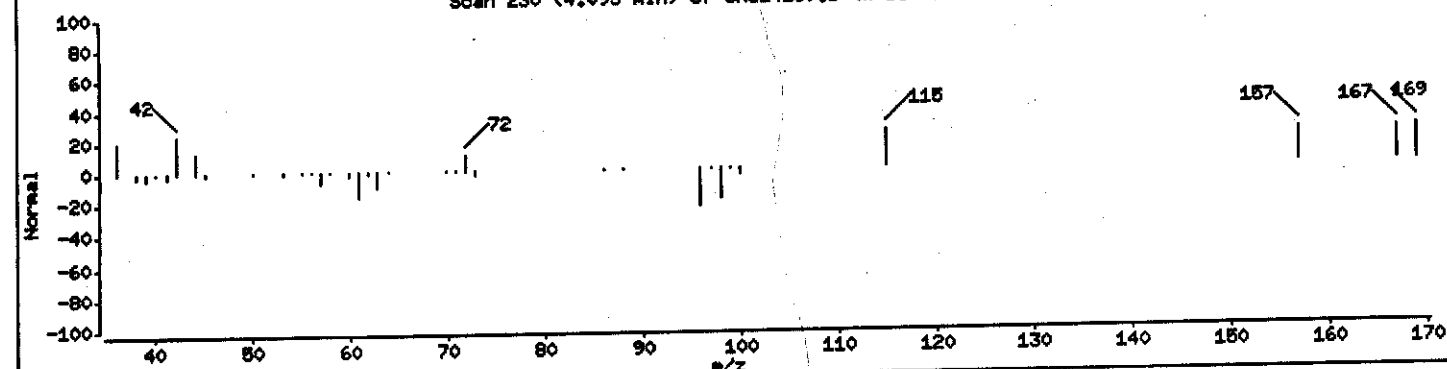
Scan 230 (4.095 min) of UXJ24289.D (Subtracted)



30 2-Butanone (Reference Spectrum)



Scan 230 (4.095 min) of UXJ24289.D (% DIFFERENCE)



Data File: \\qcanch04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24289.D

Date : 01-OCT-2004 12:51

Client ID: TRIP BLANK/092804

Instrument: s3ux11.i

Sample Info: GRDOD1AA,5ML/5ML

Purge Volume: 5.0

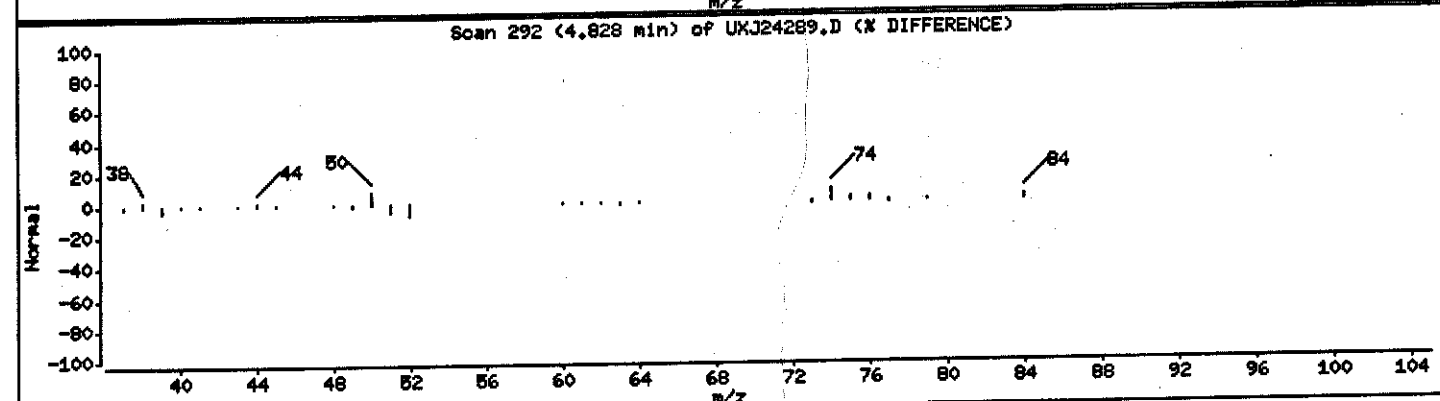
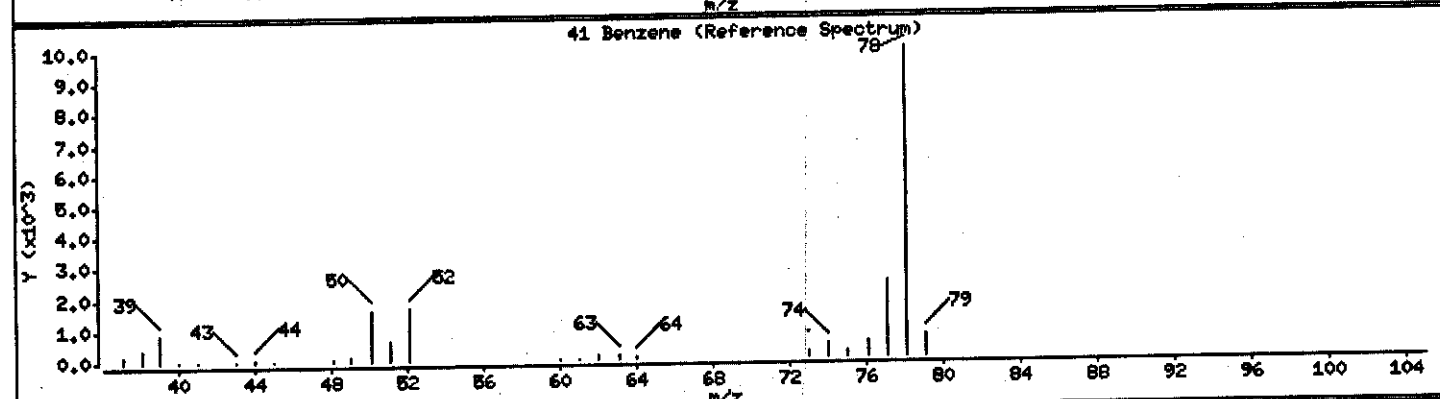
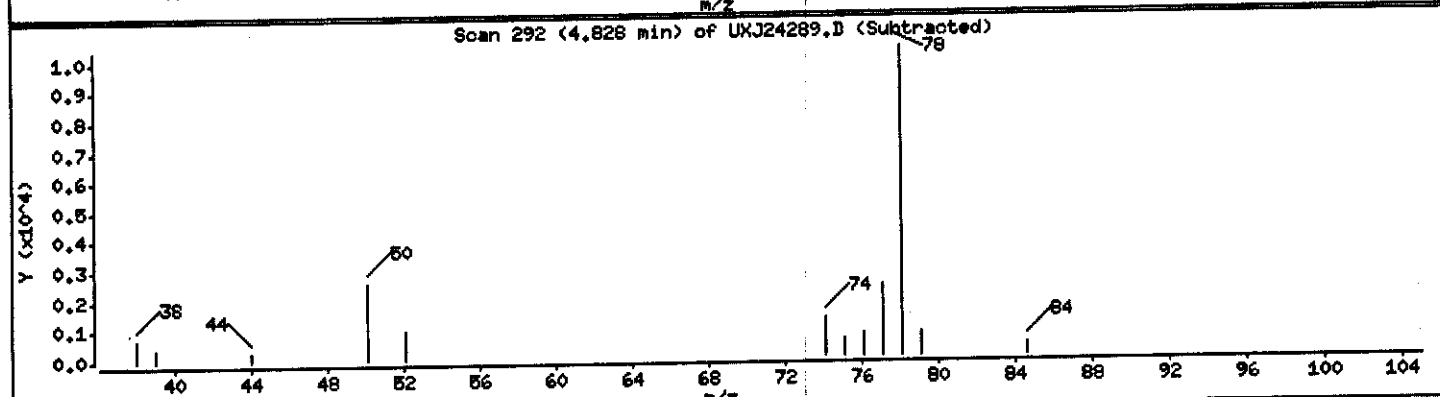
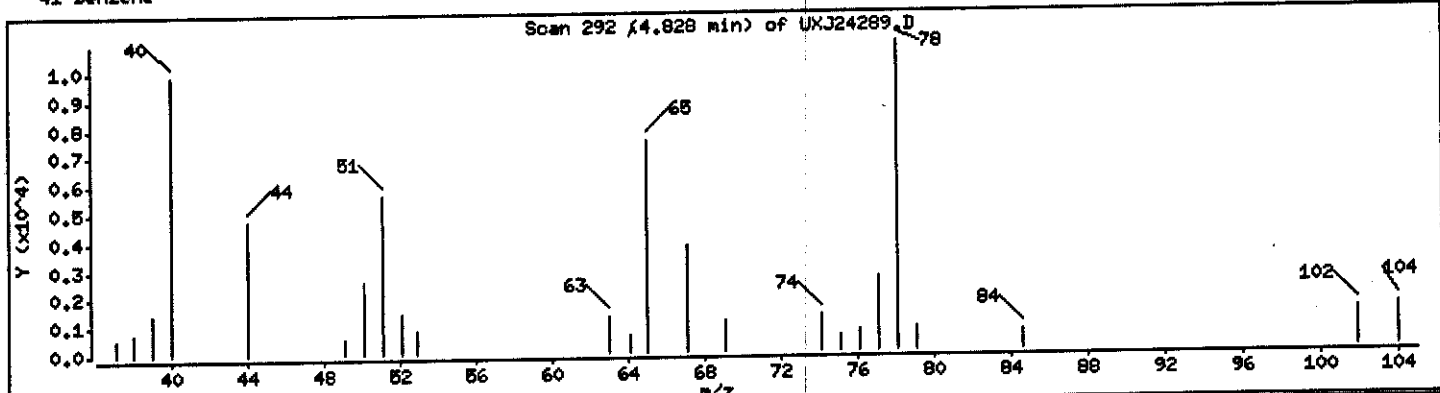
Operator: 43882

Column phase: DB624

Column diameter: 0.18

Concentration: 0.1840 ug/L

41 Benzene



Data File: \\qpanoh04\dd\chem\MSV\s3ux11.i\J41001A.b\UXJ24289.D

Date : 01-OCT-2004 12:51

Client ID: TRIP BLANK/092804

Instrument: s3ux11.i

Sample Info: GRDOD1AA,5ML/5ML

Operator: 43582

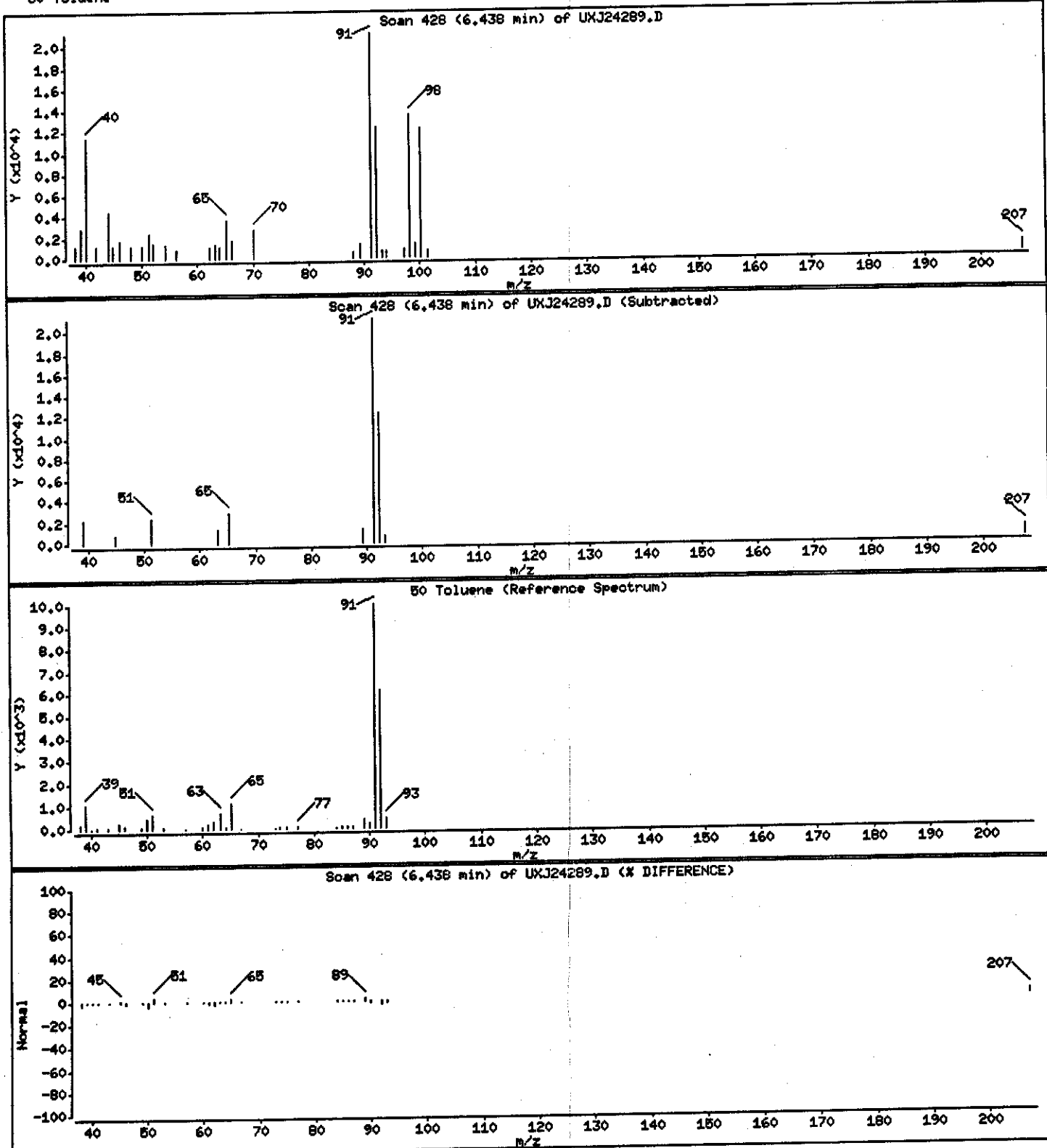
Purge Volume: 5.0

Column diameter: 0.18

Column phase: DB624

Concentration: 0.1866 ug/L

50 Toluene



SEVERN
TRENT **STL**

STANDARD DATA

Calibration History

Method : \\qcanoh04\dd\chem\MSV\aux11.i\J40914B-IC.b\8260LLUX11.m
 Start Cal Date: 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Last Cal Level: 1
 Last Cal Type : Initial Calibration

Initial Calibration

| Injection Date | Sublist | Calibration File |
|--|---------|------------------|
| Cal Level: 1 , Cal Amount: 5.000 | | |
| 14-SEP-2004 15:41 | 2-8260 | UXJ23875.D |
| 16-AUG-2004 18:11 | 3-IX | UXJ23214.D |
| Cal Level: 2 , Cal Amount: 10.000 | | |
| 14-SEP-2004 15:19 | 2-8260 | UXJ23874.D |
| 16-AUG-2004 17:48 | 3-IX | UXJ23213.D |
| Cal Level: 3 , Cal Amount: 25.000 | | |
| 14-SEP-2004 14:57 | 2-8260 | UXJ23873.D |
| 16-AUG-2004 17:26 | 3-IX | UXJ23212.D |
| Cal Level: 4 , Cal Amount: 50.000 | | |
| 14-SEP-2004 14:33 | 2-8260 | UXJ23872.D |
| 16-AUG-2004 17:03 | 3-IX | UXJ23211.D |
| Cal Level: 5 , Cal Amount: 100.00 | | |
| 14-SEP-2004 14:10 | 2-8260 | UXJ23871.D |
| 16-AUG-2004 16:40 | 3-IX | UXJ23210.D |
| Cal Level: 6 , Cal Amount: 200.00 | | |
| 14-SEP-2004 13:48 | 2-8260 | UXJ23870.D |
| 16-AUG-2004 16:18 | 3-IX | UXJ23209.D |

Continuing Calibration

| | | |
|-------------------|--------|------------|
| 14-SEP-2004 14:33 | 2-8260 | UXJ23872.D |
| | | |

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

Calibration File Names:

Level 1: \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\UXJ23875.D
 Level 2: \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\UXJ23874.D
 Level 3: \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\UXJ23873.D
 Level 4: \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\UXJ23872.D
 Level 5: \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\UXJ23871.D
 Level 6: \\qcanoh04\dd\chem\MSV\A3ux11.i\J40914B-IC.b\UXJ23870.D

| Compound | 5.000 | 10.000 | 25.000 | 50.000 | 100.000 | 200.000 | — | % RSD |
|---------------------------------|---------|---------|---------|---------|---------|---------|---------|--------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | RRF | |
| 8 Dichlorodifluoromethane | 0.28836 | 0.23754 | 0.28173 | 0.25161 | 0.24515 | 0.26401 | 0.26140 | 7.796 |
| 9 Chloromethane | 0.55674 | 0.49898 | 0.45483 | 0.43875 | 0.42154 | 0.42948 | 0.46672 | 11.132 |
| 10 Vinyl Chloride | 0.32349 | 0.31847 | 0.33246 | 0.31180 | 0.29804 | 0.33125 | 0.31925 | 4.066 |
| 11 Bromomethane | 0.17259 | 0.16446 | 0.14190 | 0.15123 | 0.13819 | 0.14020 | 0.15143 | 9.386 |
| 12 Chloroethane | 0.23244 | 0.24693 | 0.24362 | 0.23419 | 0.22089 | 0.22969 | 0.23463 | 4.043 |
| 13 Trichlorofluoromethane | 0.38407 | 0.31147 | 0.34717 | 0.32533 | 0.30602 | 0.33365 | 0.33462 | 8.499 |
| 14 Dichlorofluoromethane | 0.45871 | 0.50000 | 0.48561 | 0.49154 | 0.48286 | 0.49097 | 0.48495 | 2.915 |
| 15 Acrolein | 0.03199 | 0.03065 | 0.03129 | 0.03093 | 0.03156 | 0.03098 | 0.03123 | 1.551 |
| 16 Acetone | 0.16115 | 0.13958 | 0.11422 | 0.11384 | 0.10613 | 0.10203 | 0.12282 | 18.621 |
| 17 1,1-Dichloroethene | 0.24613 | 0.23009 | 0.19938 | 0.23265 | 0.20714 | 0.22623 | 0.22360 | 7.735 |
| 18 Freon-113 | 0.16870 | 0.16626 | 0.10968 | 0.16762 | 0.13496 | 0.15980 | 0.15117 | 15.834 |
| 19 Iodomethane | 0.31149 | 0.34851 | 0.32631 | 0.33364 | 0.33020 | 0.33284 | 0.33050 | 3.627 |
| 20 Carbon Disulfide | 0.91244 | 0.87319 | 0.74270 | 0.85398 | 0.78522 | 0.84395 | 0.83525 | 7.360 |
| 21 Methylene Chloride | 0.75227 | 0.54750 | 0.35921 | 0.31815 | 0.29068 | 0.28051 | 0.42472 | 44.272 |
| 22 Acetonitrile | 0.03290 | 0.03019 | 0.03062 | 0.02865 | 0.02976 | 0.02592 | 0.02967 | 7.789 |
| 23 Acrylonitrile | 0.09137 | 0.09573 | 0.09318 | 0.09088 | 0.09229 | 0.09022 | 0.09228 | 2.153 |
| 24 Methyl tert-butyl ether | 0.65699 | 0.67280 | 0.74332 | 0.74432 | 0.74979 | 0.73394 | 0.71686 | 5.703 |
| 25 trans-1,2-Dichloroethene | 0.28262 | 0.27489 | 0.25328 | 0.26749 | 0.25617 | 0.25710 | 0.26526 | 4.435 |
| 26 Hexane | 0.05179 | 0.04861 | 0.03324 | 0.04941 | 0.04250 | 0.04921 | 0.04579 | 15.039 |
| 27 Vinyl acetate | 0.41519 | 0.38277 | 0.42613 | 0.42027 | 0.44948 | 0.45290 | 0.42446 | 6.033 |
| 28 1,1-Dichloroethane | 0.51364 | 0.49348 | 0.46074 | 0.48352 | 0.47362 | 0.47833 | 0.48389 | 3.753 |
| 29 tert-Butyl Alcohol | 0.02052 | 0.02018 | 0.01934 | 0.01924 | 0.01980 | 0.01761 | 0.01945 | 5.262 |
| 30 2-Butanone | 0.15894 | 0.12780 | 0.13677 | 0.13376 | 0.13540 | 0.12902 | 0.13695 | 8.281 |
| M 31 1,2-Dichloroethene (total) | 0.28520 | 0.28491 | 0.26194 | 0.27101 | 0.26404 | 0.26479 | 0.27198 | 3.886 |
| 32 cis-1,2-dichloroethene | 0.28778 | 0.29492 | 0.27061 | 0.27453 | 0.27191 | 0.27248 | 0.27870 | 3.636 |

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

| Compound | 5.000 | 10.000 | 25.000 | 50.000 | 100.000 | 200.000 | RRF | % RSD |
|------------------------------|---------|---------|---------|---------|---------|---------|---------|--------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | |
| 33 2,2-Dichloropropane | 0.29593 | 0.30774 | 0.27673 | 0.29383 | 0.28018 | 0.29523 | 0.29161 | 3.907 |
| 34 Bromochloromethane | 0.12769 | 0.14200 | 0.12631 | 0.12865 | 0.12580 | 0.12388 | 0.12906 | 5.074 |
| 35 Chloroform | 0.51012 | 0.50575 | 0.47005 | 0.48650 | 0.47366 | 0.47250 | 0.48643 | 3.631 |
| 36 Tetrahydrofuran | 0.10018 | 0.08347 | 0.06844 | 0.07301 | 0.07510 | 0.07044 | 0.07844 | 15.112 |
| 37 1,1,1-Trichloroethane | 0.39287 | 0.37451 | 0.33541 | 0.38300 | 0.34846 | 0.36998 | 0.36737 | 5.878 |
| 38 1,1-Dichloropropene | 0.34921 | 0.35729 | 0.30656 | 0.35792 | 0.32912 | 0.35428 | 0.34240 | 6.001 |
| 39 Carbon Tetrachloride | 0.30286 | 0.29259 | 0.24537 | 0.31080 | 0.27649 | 0.30261 | 0.28845 | 8.383 |
| 40 1,2-Dichloroethane | 0.40025 | 0.42402 | 0.38489 | 0.39237 | 0.39169 | 0.39145 | 0.39745 | 3.498 |
| 41 Benzene | 1.25995 | 1.19011 | 1.12471 | 1.13415 | 1.09548 | 1.10254 | 1.15116 | 5.467 |
| 42 Trichloroethene | 0.27283 | 0.26390 | 0.25129 | 0.26732 | 0.25842 | 0.26258 | 0.26272 | 2.818 |
| 43 1,2-Dichloropropane | 0.29669 | 0.30122 | 0.27194 | 0.28098 | 0.27691 | 0.27607 | 0.28397 | 4.241 |
| 44 1,4-Dioxane | 0.00242 | 0.00233 | 0.00258 | 0.00261 | 0.00260 | 0.00206 | 0.00243 | 8.907 |
| 45 Dibromomethane | 0.15285 | 0.16666 | 0.15233 | 0.15822 | 0.15862 | 0.15545 | 0.15736 | 3.340 |
| 46 Bromodichloromethane | 0.38193 | 0.38487 | 0.36100 | 0.37042 | 0.36931 | 0.37381 | 0.37355 | 2.345 |
| 47 2-Chloroethyl vinyl ether | 0.13950 | 0.14798 | 0.15985 | 0.16693 | 0.17284 | 0.17092 | 0.15967 | 8.385 |
| 48 cis-1,3-Dichloropropene | 0.43417 | 0.47734 | 0.43633 | 0.45487 | 0.46634 | 0.47331 | 0.45706 | 4.058 |
| 49 4-Methyl-2-pentanone | 0.23661 | 0.23635 | 0.24230 | 0.25213 | 0.25538 | 0.25182 | 0.24577 | 3.424 |
| 50 Toluene | 1.48717 | 1.54897 | 1.47049 | 1.49192 | 1.46610 | 1.49497 | 1.49327 | 1.987 |
| 51 trans-1,3-Dichloropropene | 0.50647 | 0.53118 | 0.51429 | 0.54236 | 0.55606 | 0.56198 | 0.53539 | 4.162 |
| 52 Ethyl Methacrylate | 0.38934 | 0.45025 | 0.45000 | 0.48061 | 0.49453 | 0.49921 | 0.46066 | 8.865 |
| 53 1,1,2-Trichloroethane | 0.30188 | 0.31382 | 0.29748 | 0.30352 | 0.30279 | 0.29789 | 0.30290 | 1.953 |
| 54 1,3-Dichloropropane | 0.57206 | 0.59182 | 0.56392 | 0.55975 | 0.57182 | 0.56489 | 0.57071 | 1.996 |
| 55 Tetrachloroethene | 0.25509 | 0.26206 | 0.22142 | 0.24509 | 0.22718 | 0.24073 | 0.24193 | 6.478 |
| 56 2-Hexanone | 0.22234 | 0.23807 | 0.24231 | 0.24889 | 0.24606 | 0.25308 | 0.24179 | 4.486 |
| 57 Dibromochloromethane | 0.30967 | 0.33372 | 0.31997 | 0.31951 | 0.32624 | 0.32930 | 0.32307 | 2.641 |
| 58 1,2-Dibromoethane | 0.26921 | 0.31637 | 0.29041 | 0.30230 | 0.30732 | 0.30501 | 0.29844 | 5.561 |
| 59 Chlorobenzene | 0.96525 | 1.03980 | 0.94396 | 0.94706 | 0.94928 | 0.95592 | 0.96688 | 3.777 |
| 60 1,1,1,2-Tetrachloroethane | 0.33779 | 0.34414 | 0.32638 | 0.32925 | 0.34031 | 0.33757 | 0.33590 | 2.013 |
| 61 Ethylbenzene | 0.46024 | 0.50768 | 0.46972 | 0.48906 | 0.48935 | 0.51358 | 0.48827 | 4.245 |
| 62 m + p-Xylene | 0.60709 | 0.64675 | 0.61837 | 0.63926 | 0.62027 | 0.64155 | 0.62888 | 2.511 |
| M 63 Xylenes (total) | 0.59346 | 0.64358 | 0.61984 | 0.63276 | 0.62064 | 0.64001 | 0.62505 | 2.925 |
| 64 Xylene-o | 0.56621 | 0.63725 | 0.62277 | 0.61975 | 0.62137 | 0.63694 | 0.61738 | 4.252 |
| 65 Styrene | 1.00309 | 1.11779 | 1.08299 | 1.11462 | 1.13838 | 1.16736 | 1.10404 | 5.143 |

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

| Compound | 5.000 | 10.000 | 25.000 | 50.000 | 100.000 | 200.000 | RRF | % RSD |
|--------------------------------|---------|---------|---------|---------|---------|---------|---------|--------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | |
| 66 Bromoform | 0.20132 | 0.22491 | 0.21511 | 0.21840 | 0.22531 | 0.23254 | 0.21960 | 4.920 |
| 67 Isopropylbenzene | 1.34498 | 1.36235 | 1.30493 | 1.41191 | 1.35691 | 1.46909 | 1.37503 | 4.180 |
| 68 1,1,2-Tetrachloroethane | 0.74074 | 0.78253 | 0.73952 | 0.75703 | 0.76116 | 0.74128 | 0.75371 | 2.237 |
| 69 1,4-Dichloro-2-butene | 0.23161 | 0.23946 | 0.23911 | 0.25172 | 0.26183 | 0.26419 | 0.24799 | 5.373 |
| 70 1,2,3-Trichloropropane | 0.25299 | 0.25205 | 0.24157 | 0.24830 | 0.24773 | 0.24135 | 0.24733 | 2.016 |
| 71 Bromobenzene | 0.74497 | 0.77908 | 0.72155 | 0.75186 | 0.74513 | 0.73336 | 0.74599 | 2.605 |
| 72 n-Propylbenzene | 0.71725 | 0.70753 | 0.62905 | 0.72290 | 0.67589 | 0.72046 | 0.69551 | 5.299 |
| 73 2-Chlorotoluene | 0.71622 | 0.69234 | 0.65887 | 0.70074 | 0.67558 | 0.68895 | 0.68879 | 2.887 |
| 74 1,3,5-Trimethylbenzene | 2.25968 | 2.26774 | 2.20534 | 2.39411 | 2.31559 | 2.44388 | 2.31439 | 3.874 |
| 75 4-Chlorotoluene | 0.76654 | 0.75820 | 0.70598 | 0.74482 | 0.71695 | 0.72231 | 0.73580 | 3.304 |
| 76 tert-Butylbenzene | 1.89049 | 1.94223 | 1.70074 | 1.94123 | 1.83516 | 1.94010 | 1.87499 | 5.079 |
| 77 1,2,4-Trimethylbenzene | 2.33163 | 2.49724 | 2.33803 | 2.57507 | 2.50096 | 2.58086 | 2.47063 | 4.492 |
| 78 sec-Butylbenzene | 2.66025 | 2.55624 | 2.28546 | 2.65769 | 2.44749 | 2.62040 | 2.53792 | 5.801 |
| 79 4-Isopropyltoluene | 2.02816 | 2.10772 | 1.90515 | 2.27140 | 2.10588 | 2.25071 | 2.11151 | 6.511 |
| 80 1,3-Dichlorobenzene | 1.50394 | 1.41628 | 1.30596 | 1.36727 | 1.33637 | 1.33353 | 1.37723 | 5.270 |
| 81 1,4-Dichlorobenzene | 1.49216 | 1.50657 | 1.37893 | 1.45011 | 1.40876 | 1.40879 | 1.44089 | 3.529 |
| 82 n-Butylbenzene | 1.86201 | 1.92154 | 1.64744 | 2.01989 | 1.84399 | 2.02271 | 1.88626 | 7.389 |
| 83 1,2-Dichlorobenzene | 1.44948 | 1.40634 | 1.30068 | 1.35160 | 1.32364 | 1.30712 | 1.35648 | 4.397 |
| 84 1,2-Dibromo-3-chloropropane | 0.12641 | 0.12593 | 0.12612 | 0.12831 | 0.13070 | 0.12832 | 0.12763 | 1.446 |
| 85 1,2,4-Trichlorobenzene | 0.56391 | 0.60976 | 0.52246 | 0.58135 | 0.57921 | 0.60830 | 0.57750 | 5.596 |
| 86 Hexachlorobutadiene | 0.39076 | 0.31635 | 0.25219 | 0.27519 | 0.23759 | 0.23788 | 0.28500 | 20.941 |
| 87 Naphthalene | 1.19572 | 1.26888 | 1.21865 | 1.39469 | 1.53842 | 1.56161 | 1.36300 | 11.780 |
| 88 1,2,3-Trichlorobenzene | 0.38318 | 0.44192 | 0.36462 | 0.41931 | 0.44173 | 0.43401 | 0.41413 | 7.910 |
| 89 Ethyl Ether | 0.26862 | 0.25572 | 0.24190 | 0.23691 | 0.23958 | 0.23653 | 0.24654 | 5.241 |
| 90 Ethanol | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | <- |
| 91 3-Chloropropene | 0.08501 | 0.09752 | 0.10420 | 0.10696 | 0.10892 | 0.11570 | 0.10305 | 10.335 |
| 92 Isopropyl Ether | 0.19894 | 0.21144 | 0.22424 | 0.23464 | 0.23633 | 0.23561 | 0.22353 | 6.886 |
| 93 2-Chloro-1,3-butadiene | 0.33010 | 0.35251 | 0.38261 | 0.39081 | 0.38671 | 0.39380 | 0.37276 | 6.882 |
| 94 Propionitrile | 0.04383 | 0.04396 | 0.04410 | 0.04111 | 0.04094 | 0.03990 | 0.04231 | 4.404 |
| 95 Ethyl Acetate | 0.25153 | 0.23918 | 0.24678 | 0.23828 | 0.24450 | 0.25022 | 0.24508 | 2.253 |
| 96 Methacrylonitrile | 0.15470 | 0.15983 | 0.15530 | 0.15862 | 0.16298 | 0.16195 | 0.15890 | 2.134 |
| 97 Isobutanol | 0.01214 | 0.01067 | 0.01151 | 0.01109 | 0.01159 | 0.01152 | 0.01142 | 4.362 |
| 98 Cyclohexane | 0.37486 | 0.38746 | 0.29089 | 0.40866 | 0.34684 | 0.39833 | 0.36784 | 11.783 |

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

| Compound | 5.000 | 10.000 | 25.000 | 50.000 | 100.000 | 200.000 | — | % RSD |
|----------------------------------|---------|---------|---------|---------|---------|---------|---------|------------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | RRF | |
| 99 n-Butanol | 0.00708 | 0.00755 | 0.00826 | 0.00839 | 0.00871 | 0.00930 | 0.00822 | 9.712 <- |
| 100 Methyl Methacrylate | 0.17898 | 0.17553 | 0.18573 | 0.19952 | 0.20913 | 0.22296 | 0.19531 | 9.504 |
| 101 2-Nitropropane | 0.05322 | 0.06193 | 0.06220 | 0.06218 | 0.06202 | 0.06322 | 0.06079 | 6.151 |
| 102 Chloropicrin | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 103 Cyclohexanone | 0.02167 | 0.02329 | 0.02670 | 0.02927 | 0.03085 | 0.03125 | 0.02717 | 14.734 |
| 104 Pentachloroethane | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 105 Benzyl Chloride | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 134 Thiophene | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 135 Crotononitrile(1st Isomer) | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 136 Crotononitrile(2nd Isomer) | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| M 137 Total Crotononitrile | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 138 Paraldehyde | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 139 3,3,5-Trimethylcyclohexanone | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 140 1-Chlorohexane | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 141 1,3,5-Trichlorobenzene | 0.85746 | 0.79676 | 0.67661 | 0.75898 | 0.73463 | 0.74943 | 0.76231 | 7.986 |
| 143 Methyl Acetate | 0.22313 | 0.21756 | 0.20107 | 0.19563 | 0.19918 | 0.19750 | 0.20568 | 5.658 |
| 144 Methylcyclohexane | 0.34643 | 0.31604 | 0.23920 | 0.34078 | 0.28293 | 0.32667 | 0.30868 | 13.218 |
| 145 Dimethoxymethane | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| 146 2-Methylnaphthalene | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ <- |
| \$ 4 Dibromofluoromethane | 0.22899 | 0.23170 | 0.22899 | 0.23960 | 0.23428 | 0.23320 | 0.23279 | 1.705 |
| \$ 5 1,2-Dichloroethane-d4 | 0.31268 | 0.30905 | 0.31321 | 0.31488 | 0.33897 | 0.33187 | 0.32011 | 3.817 |
| \$ 6 Toluene-d8 | 1.15061 | 1.17656 | 1.24969 | 1.21381 | 1.20619 | 1.21032 | 1.20120 | 2.832 |
| \$ 7 Bromofluorobenzene | 0.48378 | 0.51078 | 0.51438 | 0.50777 | 0.52150 | 0.52405 | 0.51038 | 2.827 |

STL North Canton

INITIAL CALIBRATION DATA

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;start Cal Date   : 16-AUG-2004 16:18
;End Cal Date    : 14-SEP-2004 15:41
;Quant Method    : ISTD
;Target Version  : 4.04
;Integrator      :
;Method file     : HP RTE
;Cal Date        : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D
;                         14-Sep-2004 16:57 tapsvc

```

Calibration File Names:

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level 1: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D
level 2: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D
level 3: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D
level 4: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D
level 5: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D
level 6: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ2387.D

```

| Compound | 5.0000 | 10.0000 | 25.0000 | 50.0000 | 100.0000 | 200.0000 | Curve | b | Coefficients | RSD | or R^2 |
|---------------------------|---------|---------|---------|---------|----------|--------------|-------|----------|--------------|----------|---------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | ml | m2 | | |
| 8 Dichlorodifluoromethane | 0.28836 | 0.23754 | 0.28173 | 0.25161 | 0.24515 | 0.26401 AVRG | | 0.28140 | | 7.70645 | |
| 9 Chloroethylene | 0.55674 | 0.49998 | 0.45483 | 0.43875 | 0.42154 | 0.42948 AVRG | | 0.46672 | | 11.13244 | |
| 10 Vinyl Chloride | 0.32349 | 0.31447 | 0.33246 | 0.31180 | 0.29804 | 0.31125 AVRG | | 0.31925 | | 4.06579 | |
| 11 Bromomethane | 0.17259 | 0.16446 | 0.14190 | 0.15123 | 0.13819 | 0.14020 AVRG | | 0.15143 | | 9.38575 | |
| 12 Chloroethane | 0.23244 | 0.24693 | 0.24362 | 0.23419 | 0.22089 | 0.22869 AVRG | | 0.22463 | | 4.04335 | |
| 13 Trichlorofluoromethane | 0.38407 | 0.31147 | 0.34717 | 0.32533 | 0.30602 | 0.33165 AVRG | | 0.33462 | | 8.49945 | |
| 14 Dichlorofluoromethane | 0.45871 | 0.50000 | 0.48561 | 0.49154 | 0.48285 | 0.49097 AVRG | | 0.46495 | | 2.91496 | |
| 15 Acrolein | 0.03199 | 0.03065 | 0.03129 | 0.03093 | 0.03156 | 0.03098 AVRG | | 0.03123 | | 1.5547 | |
| 16 Acetone | 72580 | 128042 | 262942 | 530389 | 973885 | 1907733 QUAD | | -0.10527 | 9.26837 | 0.81620 | 0.99990 |

STL North Canton

INITIAL CALIBRATION DATA

```

start Cal Date : 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Quant Method : ISTD
Target Version : 4.04
Integrator : HP RTE
Method file : \\qcano04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
.cal Date : 14-Sep-2004 16:57 tapsvc

```

| Compound | 5.000 | 10.0000 | 25.0000 | 50.0000 | 100.0000 | 200.0000 | Curve | b ML | Coefficients ML | %RSD or R^2 |
|---------------------------------|---------|---------|----------|---------|----------|--------------|----------|----------|--------------------|----------------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | | | |
| 17 1,1-Dichloroethene | 0.24613 | 0.23009 | 0.19938 | 0.23265 | 0.20714 | 0.22623 AVRG | | 0.22360 | | 7.73538 |
| 18 Freon-113 | 37591 | 76261 | 126246 | 390502 | 619208 | 149012 QUAD | -0.02165 | 7.57113 | -1.97871 | 0.99415 |
| 19 Iodomethane | 0.31149 | 0.34851 | 0.32631 | 0.33364 | 0.33020 | 0.3284 AVRG | | 0.33050 | | 3.62742 |
| 20 Carbon Disulfide | 0.91244 | 0.87319 | 0.74270 | 0.85398 | 0.78522 | 0.84395 AVRG | | 0.83525 | | 7.35951 |
| 21 Methylene Chloride | 169409 | 251128 | 413458 | 741159 | 1333682 | 2622600 QUAD | -0.19662 | 3.80728 | -0.05949 | 0.99993 |
| 22 Acetonitrile | 0.03280 | 0.03019 | 0.03062 | 0.02865 | 0.02976 | 0.02592 AVRG | | 0.02967 | | 7.78866 |
| 23 Acrylonitrile | 0.09137 | 0.09573 | 0.09318 | 0.09088 | 0.09229 | 0.09022 AVRG | | 0.09228 | | 2.35289 |
| 24 Methyl tert-butyl ether | 0.65639 | 0.67280 | 0.74332 | 0.74432 | 0.74979 | 0.73394 AVRG | | 0.71686 | | 5.70257 |
| 25 trans-1,2-Dichloroethene | 0.28282 | 0.27489 | 0.25328 | 0.26749 | 0.25617 | 0.25710 AVRG | | 0.26526 | | 4.43506 |
| 26 Hexane | 11663 | 22298 | 38256 | 115096 | 195007 | 460062 QUAD | -0.00997 | 24.38425 | -20.19946 | 0.93631 |
| 27 Vinyl acetate | 0.41519 | 0.38277 | 0.42613 | 0.42027 | 0.44948 | 0.45290 AVRG | | 0.42446 | | 6.03292 |
| 28 1,1-Dichloroethane | 0.51364 | 0.49348 | 0.46074 | 0.48352 | 0.47362 | 0.47833 AVRG | | 0.48389 | | 3.75339 |
| 29 tert-Butyl Alcohol | 0.02052 | 0.02018 | 0.01934 | 0.01924 | 0.01980 | 0.01761 AVRG | | 0.01945 | | 5.28222 |
| 30 2-Butanone | 0.15894 | 0.12780 | 0.13677 | 0.13376 | 0.13540 | 0.12802 AVRG | | 0.13695 | | 8.28054 |
| M 31 1,2-Dichloroethene (total) | 0.28520 | 0.28491 | 0.263194 | 0.27101 | 0.26404 | 0.26479 AVRG | | 0.27198 | | 3.88557 |
| 32 cis-1,2-dichloroethene | 0.28778 | 0.29492 | 0.27061 | 0.27453 | 0.27191 | 0.27248 AVRG | | 0.27870 | | 3.63568 |
| 33 2,2-Dichloropropane | 0.29593 | 0.30774 | 0.27673 | 0.29383 | 0.28018 | 0.29523 AVRG | | 0.29161 | | 3.90723 |

STL North Canton

INITIAL CALIBRATION DATA

```

;start Cal Date : 16-AUG-2004 16:18
;End Cal Date : 14-SEP-2004 15:41
;Want Method : ISTD
;Target Version : 4.04
;Integrator Method file : HP RTE
;Method file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
;All Date : 14-Sep-2004 16:57 tapsvc

```

| Compound | Level 1 | 10.0000 | 25.0000 | 50.0000 | 100.0000 | 200.0000 | Curve | b | Coefficients | m1 | m2 | sRSD | or R^2 |
|------------------------------|---------|---------|---------|---------|----------|----------|-------|---------|--------------|---------|---------|---------|--------|
| 34 Bromochloromethane | 0.12769 | 0.14200 | 0.12631 | 0.12865 | 0.12580 | 0.12388 | AVRG | | 0.12905 | | | 5.07449 | |
| 35 Chloroform | 0.51012 | 0.50575 | 0.47005 | 0.48650 | 0.47366 | 0.47250 | AVRG | | 0.48643 | | | 3.63104 | |
| 36 Tetrahydrofuran | 22560 | 38288 | 78377 | 17007 | 34505 | 658534 | QARD | 0.00447 | 12.83982 | 4.66039 | 0.99940 | | |
| 37 1,1,1-Trichloroethane | 0.39287 | 0.37451 | 0.33541 | 0.38300 | 0.34846 | 0.36998 | AVRG | | 0.36737 | | | 5.87801 | |
| 38 1,1-Bichloropropene | 0.34921 | 0.35729 | 0.30656 | 0.35792 | 0.32912 | 0.35428 | AVRG | | 0.34240 | | | 6.00098 | |
| 39 Carbon Tetrachloride | 0.30286 | 0.29259 | 0.24537 | 0.31080 | 0.27649 | 0.30261 | AVRG | | 0.28845 | | | 8.38299 | |
| 40 1,2-Dichloroethane | 0.40025 | 0.42402 | 0.38489 | 0.39237 | 0.39145 | 0.39145 | AVRG | | 0.39745 | | | 3.49760 | |
| 41 Benzene | 1.25995 | 1.19011 | 1.12471 | 1.13415 | 1.09548 | 1.10254 | AVRG | | 1.15116 | | | 5.46736 | |
| 42 Trichloroethene | 0.27283 | 0.26390 | 0.25129 | 0.26732 | 0.25842 | 0.26258 | AVRG | | 0.26272 | | | 2.81819 | |
| 43 1,2-Dichloropropane | 0.29669 | 0.30122 | 0.27194 | 0.28089 | 0.27691 | 0.27607 | AVRG | | 0.28397 | | | 4.24139 | |
| 44 1,4-Dioxane | 0.09242 | 0.09233 | 0.09258 | 0.09261 | 0.09260 | 0.09206 | AVRG | | 0.09243 | | | 8.90709 | <- |
| 45 Dibromomethane | 0.15285 | 0.16666 | 0.15233 | 0.15822 | 0.15862 | 0.15545 | AVRG | | 0.15736 | | | 3.33962 | |
| 46 Bromodichloromethane | 0.38193 | 0.38487 | 0.36100 | 0.37042 | 0.36931 | 0.37381 | AVRG | | 0.37355 | | | 2.34541 | |
| 47 2-Chloroethyl vinyl ether | 0.13950 | 0.14798 | 0.15985 | 0.16683 | 0.17284 | 0.17092 | AVRG | | 0.15967 | | | 8.38480 | |
| 48 cis-1,3-Bichloropropene | 0.43347 | 0.47734 | 0.43633 | 0.45887 | 0.46634 | 0.47331 | AVRG | | 0.45706 | | | 4.05753 | |
| 49 4-Methyl-2-pentanone | 0.23661 | 0.23635 | 0.24230 | 0.25213 | 0.25538 | 0.25182 | AVRG | | 0.24577 | | | 3.42375 | |
| 50 Toluene | 1.48717 | 1.54897 | 1.47049 | 1.49192 | 1.46610 | 1.49497 | AVRG | | 1.49327 | | | 1.98694 | |

STL North Canton

INITIAL CALIBRATION DATA

```

start Cal Date : 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Want Method : ISTD
Target Version : 4.04
Integrator : HP RTE
Method file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Cal Date : 14-Sep-2004 16:57 tapsvc

```

| Compound | 5.0000 | 10.0000 | 25.0000 | 50.0000 | 100.0000 | 200.0000 | Curve | b | Coefficients | %RSD | or R^2 |
|------------------------------|---------|---------|---------|---------|----------|---------------|-------|---------|--------------|---------|--------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | m1 | m2 | | |
| 51 trans-1,3-Dichloropropene | 0.50647 | 0.53118 | 0.51429 | 0.54236 | 0.55606 | 0.56198 AVRG | | 0.52539 | | 4.16176 | |
| 52 Ethyl Methacrylate | 0.38934 | 0.45025 | 0.45000 | 0.48061 | 0.49453 | 0.49921 AVRG | | 0.40666 | | 8.86545 | |
| 53 1,1,2-Trichloroethane | 0.30181 | 0.31382 | 0.29748 | 0.30352 | 0.30279 | 0.29789 AVRG | | 0.30290 | | 1.95346 | |
| 54 1,3-Dichloropropane | 0.57206 | 0.59182 | 0.55392 | 0.55975 | 0.57182 | 0.56489 AVRG | | 0.57071 | | 1.93579 | |
| 55 Tetrachloroethylene | 0.25509 | 0.26206 | 0.22142 | 0.24509 | 0.22718 | 0.24073 AVRG | | 0.24193 | | 6.47809 | |
| 56 2-Hexanone | 0.22234 | 0.23807 | 0.22231 | 0.24889 | 0.24606 | 0.25308 AVRG | | 0.24179 | | 4.48622 | |
| 57 Dibromochloromethane | 0.30967 | 0.33372 | 0.31997 | 0.31951 | 0.32624 | 0.32330 AVRG | | 0.32307 | | 2.64150 | |
| 58 1,2-Dibromoethane | 0.26921 | 0.31637 | 0.28041 | 0.30230 | 0.30732 | 0.30501 AVRG | | 0.25844 | | 5.52079 | |
| 59 Chlorobenzene | 0.96525 | 1.03880 | 0.94396 | 0.94706 | 0.94928 | 0.95592 AVRG | | 0.95688 | | 3.77675 | |
| 60 1,1,1,2-Tetrachloroethane | 0.33779 | 0.34414 | 0.32638 | 0.32925 | 0.34031 | 0.33757 AVRG | | 0.33590 | | 2.01341 | |
| 61 Ethylbenzene | 0.46024 | 0.50768 | 0.45972 | 0.48906 | 0.48935 | 0.5158 AVRG | | 0.48827 | | 4.24550 | |
| 62 m + p-Xylene | 0.60709 | 0.64675 | 0.61837 | 0.63926 | 0.62027 | 0.64155 AVRG | | 0.62888 | | 2.53086 | |
| M 63 Xylenes (total) | 0.59346 | 0.64358 | 0.61984 | 0.63276 | 0.62064 | 0.64001 AVRG | | 0.62505 | | 2.92467 | |
| 64 Xylene-o | 0.56621 | 0.63725 | 0.62277 | 0.61975 | 0.62137 | 0.63694 AVRG | | 0.61738 | | 4.25246 | |
| 65 Styrene | 1.00309 | 1.11779 | 1.08299 | 1.11462 | 1.13838 | 1.16736 AVRG | | 1.10404 | | 5.14327 | |
| 66 Bromoform | 0.20132 | 0.22491 | 0.21511 | 0.22531 | 0.23254 | 0.23501 AVRG | | 0.21960 | | 4.91986 | |
| 67 Isopropylbenzene | 1.34498 | 1.36235 | 1.30493 | 1.41191 | 1.35691 | 1.46909 AVRG | | 1.37503 | | 4.17959 | |

STL North Canton

INITIAL CALIBRATION DATA

```

start Cal Date : 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Quant Method : ISTD
Target Version : 4.04
Integrator : HP RTE
Method file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Cal Date : 14-Sep-2004 16:57 tapsvc

```

| Compound | 5.000 | 10.000 | 25.000 | 50.000 | 100.000 | 200.000 | Curve | b | Coefficients | n1 | n2 | *RSD or R^2 |
|--------------------------------|---------|---------|---------|---------|---------|--------------|-------|---------|--------------|---------|----|----------------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | | | | | |
| 68 1,1,2,2-Tetrachloroethane | 0.74074 | 0.78253 | 0.73952 | 0.75703 | 0.76116 | 0.74128 AVRG | | 0.75371 | | 2.23695 | | |
| 69 1,4-Dichloro-2-butene | 0.23161 | 0.23946 | 0.23911 | 0.25172 | 0.26183 | 0.26119 AVRG | | 0.24799 | | 5.37306 | | |
| 70 1,2,3-Trichloropropane | 0.25299 | 0.25205 | 0.24157 | 0.24830 | 0.24773 | 0.24135 AVRG | | 0.24733 | | 2.01646 | | |
| 71 Bromobenzene | 0.74497 | 0.77908 | 0.72155 | 0.75186 | 0.74513 | 0.73336 AVRG | | 0.74599 | | 2.60484 | | |
| 72 n-Propylbenzene | 0.71725 | 0.70753 | 0.62905 | 0.72290 | 0.67589 | 0.72046 AVRG | | 0.69551 | | 5.29928 | | |
| 73 2-Chlorotoluene | 0.71622 | 0.69234 | 0.65887 | 0.70074 | 0.67558 | 0.68895 AVRG | | 0.68879 | | 2.88741 | | |
| 74 1,3,5-Trimethylbenzene | 2.2568 | 2.26774 | 2.20534 | 2.39411 | 2.3159 | 2.44388 AVRG | | 2.31439 | | 3.87363 | | |
| 75 4-Chlorotoluene | 0.76654 | 0.75820 | 0.70598 | 0.74482 | 0.71655 | 0.72231 AVRG | | 0.73580 | | 3.30376 | | |
| 76 tert-Butylbenzene | 1.89049 | 1.94223 | 1.70074 | 1.94123 | 1.83526 | 1.94010 AVRG | | 1.87499 | | 5.07853 | | |
| 77 1,2,4-Trimethylbenzene | 2.33163 | 2.49724 | 2.33803 | 2.57507 | 2.50096 | 2.58086 AVRG | | 2.47063 | | 4.49215 | | |
| 78 sec-Butylbenzene | 2.66025 | 2.55924 | 2.28546 | 2.65769 | 2.44749 | 2.62040 AVRG | | 2.53792 | | 5.80056 | | |
| 79 4-Isopropyltoluene | 2.02816 | 2.10772 | 1.90515 | 2.27140 | 2.10588 | 2.25071 AVRG | | 2.11151 | | 6.51141 | | |
| 80 1,3-Dichlorobenzene | 1.50394 | 1.41628 | 1.30596 | 1.36727 | 1.33637 | 1.33153 AVRG | | 1.37723 | | 5.27000 | | |
| 81 1,4-Dichlorobenzene | 1.49216 | 1.50657 | 1.37893 | 1.45011 | 1.40876 | 1.40379 AVRG | | 1.44089 | | 3.52947 | | |
| 82 m-Butylbenzene | 1.86201 | 1.92154 | 1.64744 | 2.01989 | 1.84399 | 2.02271 AVRG | | 1.88626 | | 7.38904 | | |
| 83 1,2-Dichlorobenzene | 1.44948 | 1.40534 | 1.30068 | 1.35160 | 1.32364 | 1.30712 AVRG | | 1.35648 | | 4.39703 | | |
| 84 1,2-Dibromo-3-chloropropane | 0.12641 | 0.12593 | 0.12612 | 0.12831 | 0.13070 | 0.12832 AVRG | | 0.12763 | | 1.44611 | | |

STL North Canton

INITIAL CALIBRATION DATA

```

;start Cal Date      : 16-AUG-2004 16:18
;End Cal Date       : 14-SEP-2004 15:41
;Methodant          : ISTD
;Target Version     : 4.04
;Integrator         :
;fethod File        : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
;all Date           : 14-Sep-2004 16:57 tapsvc

```

| Compound | 5.000 | 10.000 | 25.000 | 50.000 | 100.000 | 200.000 | Curve | b | Coefficients | RMSD |
|---------------------------|---------|---------|---------|---------|---------|--------------|------------|----------|--------------|---------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | mL | R2 | or R^2 |
| 85 1,2,4-Trichlorobenzene | 0.56391 | 0.60976 | 0.52246 | 0.58135 | 0.57921 | 0.60830 AVRG | | 0.57750 | 5.59592 | |
| 86 Hexachlorobutadiene | 36648 | 62649 | 126348 | 272701 | 475139 | 979903 QUAD | -0.06586 | 4.17449 | 0.11148 | 0.99883 |
| 87 Naphthalene | 1.19572 | 1.26888 | 1.21865 | 1.39459 | 1.55842 | 1.56161 AVRG | 1.36300 | 1.177993 | | |
| 88 1,2,3-Trichlorobenzene | 0.38318 | 0.44192 | 0.36462 | 0.41931 | 0.44173 | 0.43401 AVRG | 0.41413 | | 7.90998 | |
| 89 Ethyl Ether | 0.28862 | 0.25572 | 0.24190 | 0.23691 | 0.23958 | 0.23653 AVRG | 0.24654 | | 5.24098 | |
| 90 Ethanol | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.000e+000 | | 0.000e+000 | <- |
| 91 3-Chloropropene | 0.09501 | 0.09752 | 0.10420 | 0.10696 | 0.10892 | 0.11570 AVRG | 0.10305 | 10.33497 | | |
| 92 Isopropyl Ether | 0.19894 | 0.21144 | 0.22424 | 0.23464 | 0.2633 | 0.23561 AVRG | 0.22353 | | 6.88556 | |
| 93 2-Chloro-1,3-butadiene | 0.33010 | 0.35251 | 0.38261 | 0.39081 | 0.38671 | 0.39380 AVRG | 0.37276 | | 6.88150 | |
| 94 Propionitrile | 0.04383 | 0.04396 | 0.04410 | 0.04111 | 0.04094 | 0.03990 AVRG | 0.04231 | | 4.40359 | |
| 95 Ethyl Acetate | 0.25153 | 0.23918 | 0.24678 | 0.23828 | 0.24450 | 0.25022 AVRG | 0.24508 | | 2.25277 | |
| 96 Methacrylonitrile | 0.15470 | 0.15983 | 0.15530 | 0.15862 | 0.16298 | 0.16195 AVRG | 0.15890 | | 2.13440 | |
| 97 Isobutanol | 0.01214 | 0.01067 | 0.01151 | 0.01109 | 0.01159 | 0.01152 AVRG | 0.01142 | | 4.36240 | |
| 98 Cyclohexane | 0.37486 | 0.38746 | 0.29089 | 0.40866 | 0.36684 | 0.39833 AVRG | 0.36784 | | 1.1.78329 | |
| 99 n-Butanol | 0.00708 | 0.00755 | 0.00826 | 0.00839 | 0.00871 | 0.00930 AVRG | 0.00822 | | 9.71167<- | |
| 100 Methyl Methacrylate | 0.17898 | 0.17553 | 0.18573 | 0.19952 | 0.20813 | 0.22296 AVRG | 0.19531 | | 9.50417 | |
| 101 2-Nitropropane | 0.05322 | 0.06193 | 0.06220 | 0.06202 | 0.06322 | AVRG | 0.06079 | | 6.15064 | |

STL North Canton

INITIAL CALIBRATION DATA

```

start Cal Date : 16-AUG-2004 16:18
end Cal Date : 14-SEP-2004 15:41
quant Method : ISSTD
target Version : 4.04
integrator : HP RTE
method file : \\qcano04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
cal Date : 14-Sep-2004 16:57 tapsvc

```

| Compound | 5.0000 | 10.0000 | 25.0000 | 50.0000 | 100.0000 | 200.0000 | Curve | b | Coefficients | #RSD | or R^2 |
|---------------------------------|---------|---------|---------|---------|----------|----------|-------|-------------|--------------|------|--------|
| | Level 1 | Level 2 | Level 3 | Level 4 | Level 5 | Level 6 | | | m1 | m2 | |
| 102 Chloropicrin | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 103 Cyclohexanone | 0.02167 | 0.02329 | 0.02670 | 0.02927 | 0.03085 | 0.03125 | AVRG | 0.02717 | 14.73408 | | |
| 104 Pentachloroethane | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 105 Benzyl Chloride | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 134 Thiophene | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 135 Crotononitrile(1st Isomer) | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 136 Crotononitrile(2nd Isomer) | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| M 137 Total Crotononitrile | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 138 Paraaldehyde | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 139 3,3,5-Triethylcyclohexanone | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 140 1-Chlorohexane | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 141 1,3,5-Trichlorobenzene | 0.85546 | 0.79676 | 0.67661 | 0.75898 | 0.73463 | 0.74943 | AVRG | 0.76231 | 7.98573 | | |
| 143 Methyl Acetate | 0.22313 | 0.21756 | 0.20107 | 0.19563 | 0.19918 | 0.19750 | AVRG | 0.20568 | 5.65840 | | |
| 144 Methylcyclohexane | 0.34643 | 0.31604 | 0.23920 | 0.34078 | 0.28293 | 0.32667 | AVRG | 0.30868 | 13.21820 | | |
| 145 Dimethoxymethane | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |
| 146 2-Methylnaphthalene | +++++ | +++++ | +++++ | +++++ | +++++ | +++++ | AVRG | 0.0000e+000 | 0.0000e+000 | <- | |

STL North Canton

INITIAL CALIBRATION DATA

```

;start Cal Date      : 16-AUG-2004 16:18
;Ind Cal Date       : 14-SEP-2004 15:41
;Want Method        : ISTD
;Target Version     : 4.04
;Integrator         :
;Method file        : HP RTE
;Cal Date           : \\qcanoh04\dd\chem\MSV\aux11.i\J40914B-IC.b\8260LLUX11.m
;                         14-Sep-2004 16:57 tapsvc

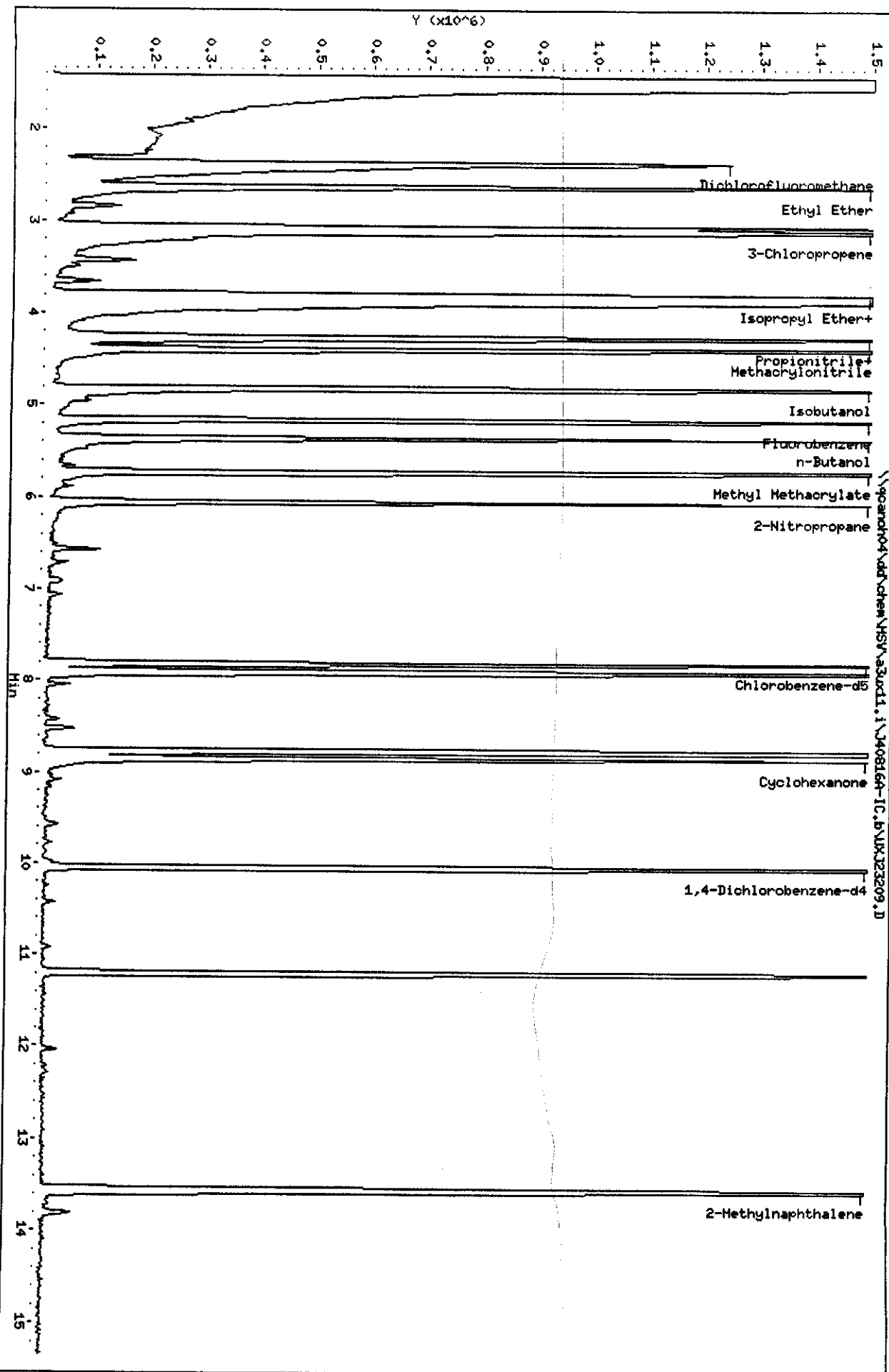
```

| Compound | 5.0000 | 10.0000 | 25.0000 | 50.0000 | 100.0000 | 200.0000 | Curve | b | Coefficients m1 | m2 | *RSD or R^2 |
|----------------------------|---------|---------|---------|---------|----------|----------|-------|---|--------------------|----|----------------|
| \$ 4 Dibromoformetane | 0.22899 | 0.23170 | 0.22899 | 0.23960 | 0.23428 | 0.23320 | AVRG | | 0.23279 | | 1.70494 |
| \$ 5 1,2-Dichloroethane-d4 | 0.31268 | 0.30905 | 0.31321 | 0.31488 | 0.33897 | 0.33187 | AVRG | | 0.32011 | | 3.81724 |
| \$ 6 Toluene-d8 | 1.15961 | 1.17656 | 1.24969 | 1.21381 | 1.20619 | 1.21032 | AVRG | | 1.20120 | | 2.83165 |
| \$ 7 Bromofluorobenzene | 0.48378 | 0.51078 | 0.51438 | 0.50777 | 0.52150 | 0.52405 | AVRG | | 0.51038 | | 2.82658 |

| Curve | Formula | Units |
|----------|-----------------------------|----------|
| Averaged | Amt = Rsp/m1 | Response |
| Quad | Amt = b + m1*Rsp + m2*Rsp^2 | Response |

Data File: \\pcando4\dd\chem\MSV\aa3dx11.i\\40816A-IC.b\\XJ23209.D
Date : 16-AUG-2004 16:18
Client ID:
Sample Info: 200NG-R91C
Purge Volume: 5.0
Column phase: DB-24

Instrument: a3dx11.i
Operator: 43602
Column diameter: 0.18
\\pcando4\dd\chem\MSV\aa3dx11.i\\40816A-IC.b\\XJ23209.D



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\UXJ23209.D
Lab Smp Id: 200NG-A9IC
Inj Date : 16-AUG-2004 16:18
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 200NG-A9IC
Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 6
Comment :
Method : \\QCANOH04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:56 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 8 Calibration Sample, Level: 6
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|----------------------------|-----------|---------|----------------|----------|---------|------------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| * 1 Fluorobenzene | 96 | 5.171 | 5.171 (1.000) | 2040512 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.822 | 7.822 (1.000) | 14444382 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 10.046 | 10.046 (1.000) | 767740 | 50.0000 | | |
| 14 Dichlorofluoromethane | 67 | 2.367 | 2.367 (0.458) | 4007314 | 200.000 | 202.48 (A) | |
| 89 Ethyl Ether | 59 | 2.627 | 2.627 (0.508) | 1930594 | 200.000 | 191.88 | |
| 91 3-Chloropropene | 76 | 3.112 | 3.112 (0.602) | 944222 | 200.000 | 224.54 (A) | |
| 92 Isopropyl Ether | 87 | 3.810 | 3.810 (0.737) | 9615196 | 1000.00 | 1054.0 (A) | |
| 93 2-Chloro-1,3-butadiene | 53 | 3.846 | 3.846 (0.744) | 3214209 | 200.000 | 211.29 (A) | |
| 94 Propionitrile | 54 | 4.260 | 4.260 (0.824) | 6513111 | 400.000 | 377.22 (A) | |
| 95 Ethyl Acetate | 43 | 4.260 | 4.260 (0.824) | 4084685 | 400.000 | 408.39 (A) | |
| 96 Methacrylonitrile | 41 | 4.390 | 4.390 (0.849) | 1321846 | 200.000 | 203.84 (A) | |
| 97 Isobutanol | 41 | 4.816 | 4.816 (0.616) | 1331366 | 4000.00 | 4036.3 (A) | |
| 99 n-Butanol | 56 | 5.361 | 5.361 (0.685) | 1074873 | 4000.00 | 4527.6 (A) | |
| 100 Methyl Methacrylate | 41 | 5.727 | 5.727 (1.108) | 1819805 | 200.000 | 228.32 (A) | |
| 101 2-Nitropropane | 41 | 6.059 | 6.059 (1.172) | 1031981 | 400.000 | 415.94 (A) | |
| 103 Cyclohexanone | 55 | 8.851 | 8.851 (0.881) | 959642 | 2000.00 | 2300.0 (A) | |
| 146 2-Methylnaphthalene | 142 | 13.561 | 13.561 (1.350) | 4079814 | 400.000 | 1085.6 (A) | |

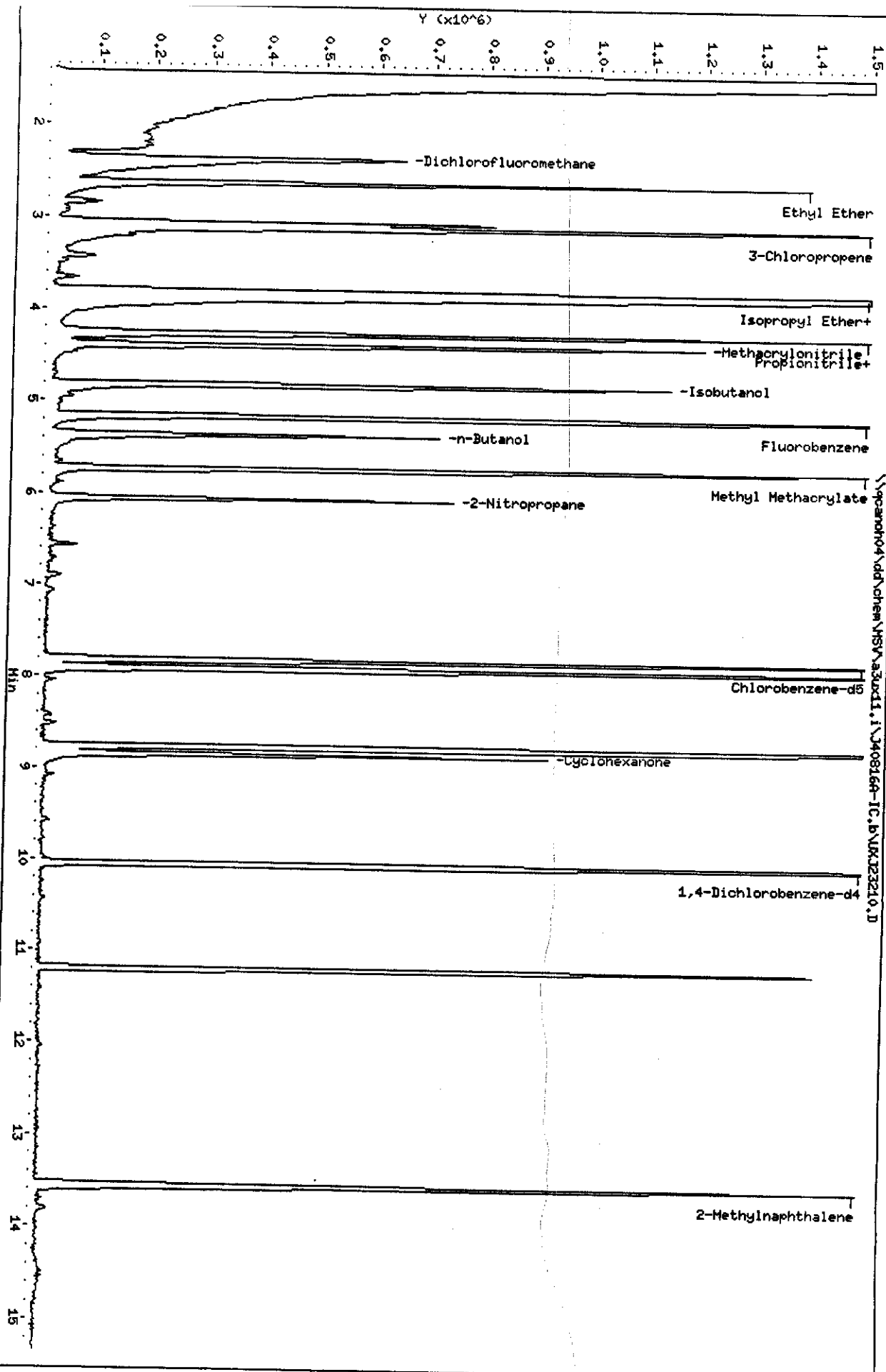
Data File: \\gcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23209.D
Report Date: 17-Aug-2004 14:56

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\pcando4\\dd\\chem\\MSV\\a30x11.i\\J40816A-IC.b\\UKJ23210.D
Date : 16-AUG-2004 16:40
Client ID:
Sample Info: 1000G-A91C
Purge Volume: 5.0
Column Phase: DB224

Instrument: a30x11.i
Operator: 435532
Column diameter: 0.18
\\pcando4\\dd\\chem\\MSV\\a30x11.i\\J40816A-IC.b\\UKJ23210.D



STL North Canton

VOLATILE REPORT SW-846 Method
 Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\UXJ23210.D
 Lab Smp Id: 100NG-A9IC
 Inj Date : 16-AUG-2004 16:40
 Operator : 43582 Inst ID: A3UX11.i
 Smp Info : 100NG-A9IC
 Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 5
 Comment :
 Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\8260LLUX11.m
 Meth Date : 17-Aug-2004 14:56 evans1 Quant Type: ISTD
 Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
 Als bottle: 9 Calibration Sample, Level: 5
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: 3-IX.SUB
 Target Version: 4.04
 Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|----------------------------|-----------|---------|----------------|---------|---------|------------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| * 1 Fluorobenzene | 96 | 5.171 | 5.171 (1.000) | 1987706 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.822 | 7.822 (1.000) | 1429041 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 10.046 | 10.046 (1.000) | 734335 | 50.0000 | | |
| 14 Dichlorofluoromethane | 67 | 2.367 | 2.367 (0.458) | 1919583 | 100.000 | 99.570 | |
| 89 Ethyl Ether | 59 | 2.627 | 2.627 (0.508) | 952411 | 100.000 | 97.174 | |
| 91 3-Chloropropene | 76 | 3.112 | 3.112 (0.602) | 432982 | 100.000 | 105.69 | |
| 92 Isopropyl Ether | 87 | 3.810 | 3.810 (0.737) | 4697492 | 500.000 | 528.62 (A) | |
| 93 2-Chloro-1,3-butadiene | 53 | 3.846 | 3.846 (0.744) | 1537316 | 100.000 | 103.74 | |
| 94 Propionitrile | 54 | 4.260 | 4.260 (0.824) | 325535 | 200.000 | 193.55 | |
| 95 Ethyl Acetate | 43 | 4.260 | 4.260 (0.824) | 1943985 | 200.000 | 199.52 | |
| 96 Methacrylonitrile | 41 | 4.390 | 4.390 (0.849) | 647907 | 100.000 | 102.57 | |
| 97 Isobutanol | 41 | 4.816 | 4.816 (0.616) | 662388 | 2000.00 | 2029.7 (A) | |
| 99 n-Butanol | 56 | 5.361 | 5.361 (0.685) | 498130 | 2000.00 | 2120.8 (A) | |
| 100 Methyl Methacrylate | 41 | 5.727 | 5.727 (1.108) | 831362 | 100.000 | 107.08 | |
| 101 2-Nitropropane | 41 | 6.059 | 6.059 (1.172) | 493098 | 200.000 | 204.02 (A) | |
| 103 Cyclohexanone | 55 | 8.851 | 8.851 (0.881) | 453132 | 1000.00 | 1135.4 (A) | |
| 146 2-Methylnaphthalene | 142 | 13.561 | 13.561 (1.350) | 1299383 | 200.000 | 361.47 (A) | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23210.D
Report Date: 17-Aug-2004 14:57

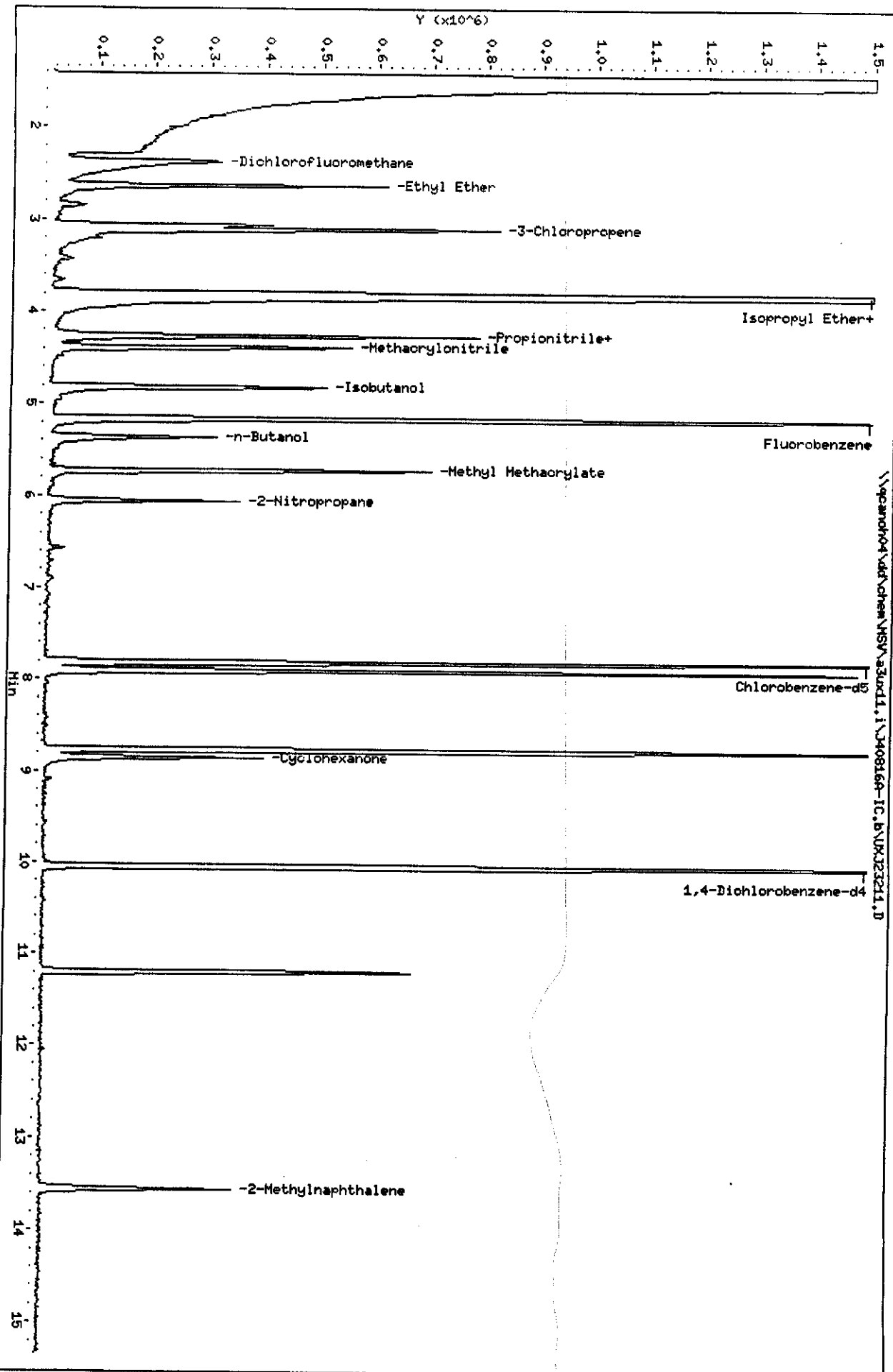
QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\pcanoh04\\dat\\chen\\HSV\\a30x11.i\\40816A-IC.b\\UKJ323214.D
Date : 16-AUG-2004 17:03
Client ID:
Sample Info: 50NG-A9IC

Purge Volume: 5.0
Column Phase: DBc24

Instrument: a30x11.i
Operator: 435622
Column diameter: 0.18



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\UXJ23211.D
Lab Smp Id: 50NG-A9IC
Inj Date : 16-AUG-2004 17:03
Operator : 43582 Inst ID: A3UX11.i
Smp Info : 50NG-A9IC
Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 4
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:57 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 10 Calibration Sample, Level: 4
Dil Factor: 1.00000
Integrator: HP RTE
Target Version: 4.04 Compound Sublist: 3-IX.SUB
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|----------------------------|-----------|---------|--------|---------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * 1 Fluorobenzene | 96 | 5.171 | 5.171 | (1.000) | 1946935 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.822 | 7.822 | (1.000) | 1394264 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 10.046 | 10.046 | (1.000) | 698314 | 50.0000 | | |
| 14 Dichlorofluoromethane | 67 | 2.379 | 2.379 | (0.460) | 957001 | 50.0000 | 50.680 | |
| 89 Ethyl Ether | 59 | 2.639 | 2.639 | (0.510) | 461257 | 50.0000 | 48.047 | |
| 91 3-Chloropropene | 76 | 3.112 | 3.112 | (0.602) | 208248 | 50.0000 | 51.898 | |
| 92 Isopropyl Ether | 87 | 3.810 | 3.810 | (0.737) | 2284128 | 250.000 | 262.42 (A) | |
| 93 2-Chloro-1,3-butadiene | 53 | 3.846 | 3.846 | (0.744) | 760882 | 50.0000 | 52.422 | |
| 94 Propionitrile | 54 | 4.260 | 4.260 | (0.824) | 160094 | 100.000 | 97.178 | |
| 95 Ethyl Acetate | 43 | 4.272 | 4.272 | (0.826) | 927819 | 100.000 | 97.223 | |
| 96 Methacrylonitrile | 41 | 4.390 | 4.390 | (0.849) | 308829 | 50.0000 | 49.914 | |
| 97 Isobutanol | 41 | 4.816 | 4.816 | (0.616) | 309158 | 1000.00 | 970.97 (A) | |
| 99 n-Butanol | 56 | 5.361 | 5.361 | (0.685) | 234037 | 1000.00 | 1021.2 (A) | |
| 100 Methyl Methacrylate | 41 | 5.727 | 5.727 | (1.108) | 388455 | 50.0000 | 51.079 | |
| 101 2-Nitropropane | 41 | 6.059 | 6.059 | (1.172) | 242106 | 100.000 | 102.27 | |
| 103 Cyclohexanone | 55 | 8.851 | 8.851 | (0.881) | 204421 | 500.000 | 538.66 (A) | |
| 146 2-Methylnaphthalene | 142 | 13.561 | 13.561 | (1.350) | 247408 | 100.000 | 72.376 | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23211.D
Report Date: 17-Aug-2004 14:57

QC Flag Legend

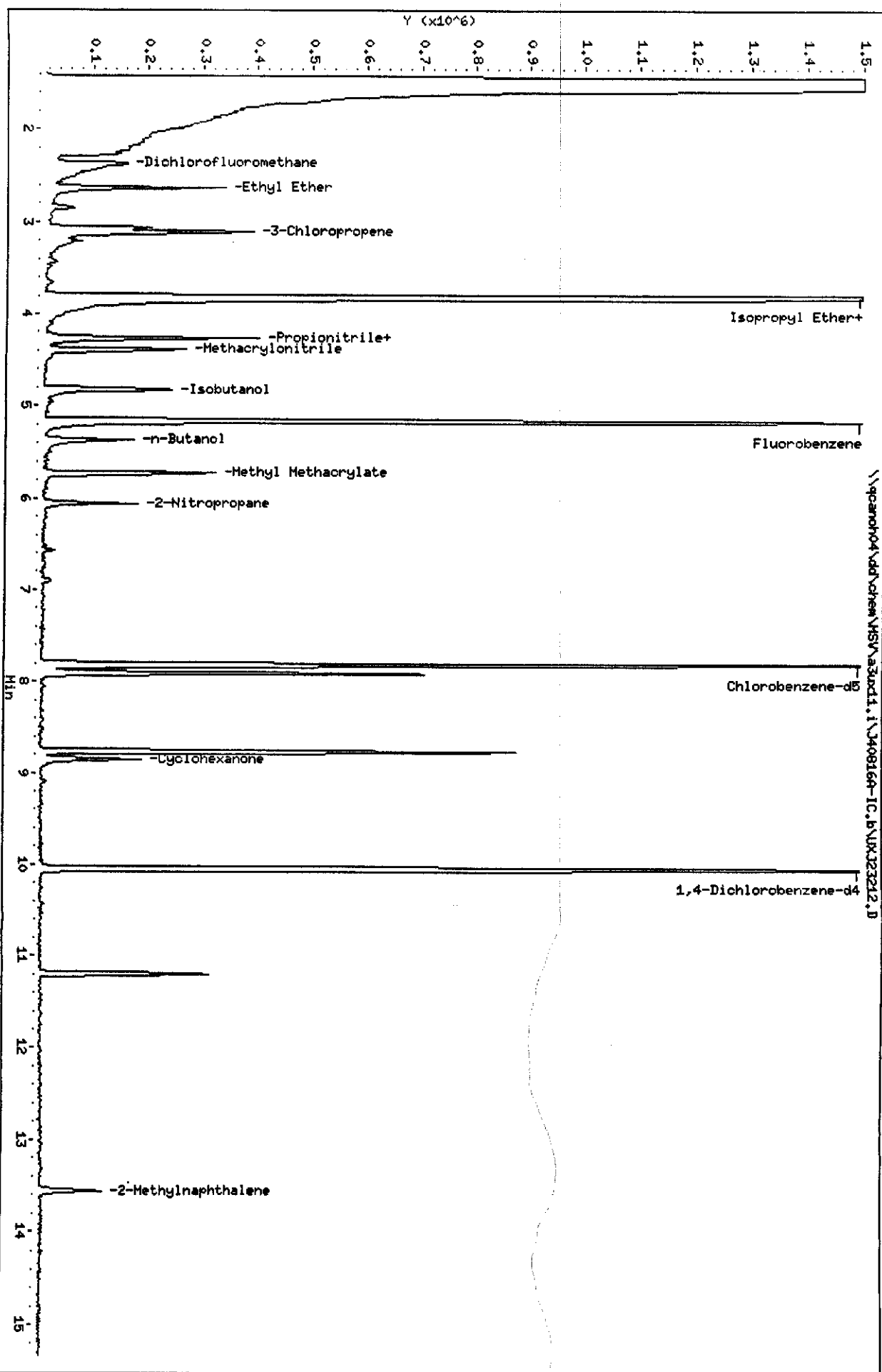
A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\qcaroh04\\d\\chem\\HSV\\a3ux11.i\\J40816A-IC.b\\UX323212.D
Date : 16-AUG-2004 17:26
Client ID:
Sample Info: 25NG-A9IC
Purge Volume: 5.0
Column phase: DB624

Instrument: a3ux11.i

Operator: 43582

Column diameter: 0.18
\\qcaroh04\\d\\chem\\HSV\\a3ux11.i\\J40816A-IC.b\\UX323212.D



STL North Canton

VOLATILE REPORT SW-846 Method
Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\UXJ23212.D
Lab Smp Id: 25NG-A9IC
Inj Date : 16-AUG-2004 17:26
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 25NG-A9IC
Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 3
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:57 evans1 Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 11 Calibration Sample, Level: 3
Dil Factor: 1.00000
Integrator: HP RTE
Target Version: 4.04 Compound Sublist: 3-IX.SUB
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|----------------------------|-----------|---------|----------------|---------|---------|-----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| * 1 Fluorobenzene | 96 | 5.171 | 5.171 (1.000) | 1894679 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.822 | 7.822 (1.000) | 1362249 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 10.046 | 10.046 (1.000) | 685784 | 50.0000 | | |
| 14 Dichlorofluoromethane | 67 | 2.378 | 2.378 (0.460) | 460035 | 25.0000 | 25.034 | |
| 89 Ethyl Ether | 59 | 2.639 | 2.639 (0.510) | 229157 | 25.0000 | 24.529 | |
| 91 3-Chloropropene | 76 | 3.112 | 3.112 (0.602) | 98714 | 25.0000 | 25.279 | |
| 92 Isopropyl Ether | 87 | 3.810 | 3.810 (0.737) | 1062158 | 125.000 | 125.40 | |
| 93 2-Chloro-1,3-butadiene | 53 | 3.846 | 3.846 (0.744) | 362465 | 25.0000 | 25.661 | |
| 94 Propionitrile | 54 | 4.260 | 4.260 (0.824) | 83549 | 50.0000 | 52.114 | |
| 95 Ethyl Acetate | 43 | 4.272 | 4.272 (0.826) | 467574 | 50.0000 | 50.347 | |
| 96 Methacrylonitrile | 41 | 4.390 | 4.390 (0.849) | 147119 | 25.0000 | 24.434 | |
| 97 Isobutanol | 41 | 4.816 | 4.816 (0.616) | 156732 | 500.000 | 503.82(A) | |
| 99 n-Butanol | 56 | 5.372 | 5.372 (0.687) | 112573 | 500.000 | 502.78(A) | |
| 100 Methyl Methacrylate | 41 | 5.727 | 5.727 (1.108) | 175947 | 25.0000 | 23.774 | |
| 101 2-Nitropropane | 41 | 6.059 | 6.059 (1.172) | 117850 | 50.0000 | 51.156 | |
| 103 Cyclohexanone | 55 | 8.851 | 8.851 (0.881) | 91550 | 250.000 | 245.65(A) | |
| 146 2-Methylnaphthalene | 142 | 13.561 | 13.561 (1.350) | 80002 | 50.0000 | 23.831 | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23212.D
Report Date: 17-Aug-2004 14:58

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\pcando4\\dd\\chem\\HSV\\a30x11.i\\J40816A-IC.b\\UX323213.D
Date : 16-AUG-2004 17:48

Client ID:

Sample Info: long-A91C

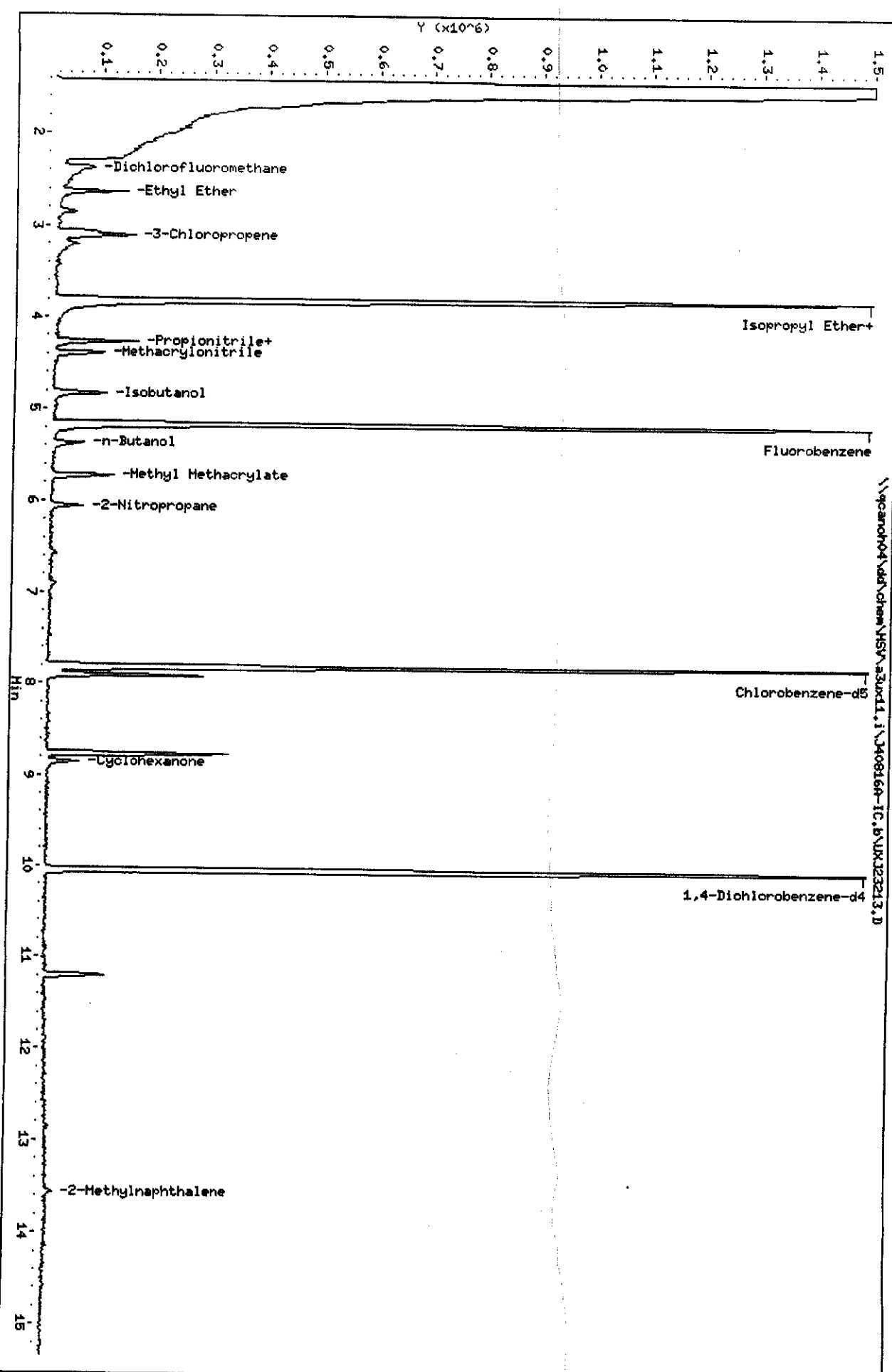
Purge Volume: 5.0

Column Phase: DB624

Instrument: a30x11.i

Operator: 43582

Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23213.D
Report Date: 17-Aug-2004 14:58

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23213.D
Lab Smp Id: 10NG-A9IC
Inj Date : 16-AUG-2004 17:48
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 10NG-A9IC
Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 2
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:58 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 12 Calibration Sample, Level: 2
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|----------------------------|-----------|---------|--------|---------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * 1 Fluorobenzene | 96 | 5.171 | 5.171 | (1.000) | 1783996 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.822 | 7.822 | (1.000) | 1312732 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 10.046 | 10.046 | (1.000) | 643754 | 50.0000 | | |
| 14 Dichlorofluoromethane | 67 | 2.379 | 2.379 | (0.460) | 178398 | 10.0000 | 10.310 | |
| 89 Ethyl Ether | 59 | 2.639 | 2.639 | (0.510) | 91239 | 10.0000 | 10.372 | |
| 91 3-Chloropropene | 76 | 3.112 | 3.112 | (0.602) | 34795 | 10.0000 | 9.463 | |
| 92 Isopropyl Ether | 87 | 3.810 | 3.810 | (0.737) | 377202 | 50.0000 | 47.294 | |
| 93 2-Chloro-1,3-butadiene | 53 | 3.846 | 3.846 | (0.744) | 125775 | 10.0000 | 9.457 | |
| 94 Propionitrile | 54 | 4.260 | 4.260 | (0.824) | 31372 | 20.0000 | 20.782 | |
| 95 Ethyl Acetate | 43 | 4.272 | 4.272 | (0.826) | 170678 | 20.0000 | 19.518 | |
| 96 Methacrylonitrile | 41 | 4.390 | 4.390 | (0.849) | 57028 | 10.0000 | 10.059 | |
| 97 Isobutanol | 41 | 4.828 | 4.828 | (0.617) | 56016 | 200.000 | 186.86 | |
| 99 n-Butanol | 56 | 5.384 | 5.384 | (0.688) | 39656 | 200.000 | 183.79 | |
| 100 Methyl Methacrylate | 41 | 5.727 | 5.727 | (1.108) | 62629 | 10.0000 | 8.987 | |
| 101 2-Nitropropane | 41 | 6.059 | 6.059 | (1.172) | 44196 | 20.0000 | 20.375 | |
| 103 Cyclohexanone | 55 | 8.851 | 8.851 | (0.881) | 29984 | 100.000 | 85.706 | |
| 146 2-Methylnaphthalene | 142 | 13.561 | 13.561 | (1.350) | 13721 | 20.0000 | 4.354 | |

Data File: \\pcancho4\\dd\\chem\\MSV\\a3\\x11.i\\J40816A-IC.b\\JK323214.D
Date : 16-AUG-2004 18:11

Client ID:
Sample Info: SNC-691C
Purge Volume: 5.0
Column Phase: DB624

Instrument: a3\\x11.i

Operator: 435882
Column diameter: 0.18

Y (x10⁶)

1.5

1.4

1.3

1.2

1.1

1.0

0.9

0.8

Y (x10⁶)

0.7

0.6

0.5

0.4

0.3

0.2

-Dichlorofluoromethane
-Ethyl Ether

-3-Chloropropene

-Isopropyl Ether+

-Propionitrile+
-Methacrylonitrile

-Isobutanol

Fluorobenzene

-n-Butanol
-Methyl Methacrylate

-2-Nitropropane

Chlorobenzene-d5

-Cyclohexanone

1,4-Dichlorobenzene-d4

-2-Methylnaphthalene

Min

8

9

10

11

12

13

14

15

Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\UXJ23214.D
Report Date: 17-Aug-2004 14:59

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\UXJ23214.D
Lab Smp Id: 5NG-A9IC
Inj Date : 16-AUG-2004 18:11
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 5NG-A9IC
Misc Info : J40816A-IC,8260LLUX11,3-IX.SUB,43582,1,1
Comment :
Method : \\QCANOH04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:59 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 13 Calibration Sample, Level: 1
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|----------------------------|-----------|---------|--------|---------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * 1 Fluorobenzene | 96 | 5.171 | 5.171 | (1.000) | 1729289 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.822 | 7.822 | (1.000) | 1295054 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 10.046 | 10.046 | (1.000) | 623017 | 50.0000 | | |
| 14 Dichlorofluoromethane | 67 | 2.379 | 2.379 | (0.460) | 79325 | 5.00000 | 4.730 | |
| 89 Ethyl Ether | 59 | 2.639 | 2.639 | (0.510) | 46453 | 5.00000 | 5.448 | |
| 91 3-Chloropropene | 76 | 3.112 | 3.112 | (0.602) | 14700 | 5.00000 | 4.124 | |
| 92 Isopropyl Ether | 87 | 3.822 | 3.822 | (0.739) | 172012 | 25.0000 | 22.250 | |
| 93 2-Chloro-1,3-butadiene | 53 | 3.846 | 3.846 | (0.744) | 57084 | 5.00000 | 4.428 | |
| 94 Propionitrile | 54 | 4.260 | 4.260 | (0.824) | 15160 | 10.0000 | 10.360 | |
| 95 Ethyl Acetate | 43 | 4.272 | 4.272 | (0.826) | 86994 | 10.0000 | 10.263 | |
| 96 Methacrylonitrile | 41 | 4.390 | 4.390 | (0.849) | 26752 | 5.00000 | 4.868 | |
| 97 Isobutanol | 41 | 4.816 | 4.816 | (0.616) | 31443 | 100.000 | 106.32 | |
| 99 n-Butanol | 56 | 5.372 | 5.372 | (0.687) | 18347 | 100.000 | 86.193 | |
| 100 Methyl Methacrylate | 41 | 5.727 | 5.727 | (1.108) | 30951 | 5.00000 | 4.582 | |
| 101 2-Nitropropane | 41 | 6.059 | 6.059 | (1.172) | 18407 | 10.0000 | 8.754 | |
| 103 Cyclohexanone | 55 | 8.851 | 8.851 | (0.881) | 13502 | 50.0000 | 39.878 | |
| 146 2-Methylnaphthalene | 142 | 13.561 | 13.561 | (1.350) | 1849 | 10.0000 | 0.6063 | |

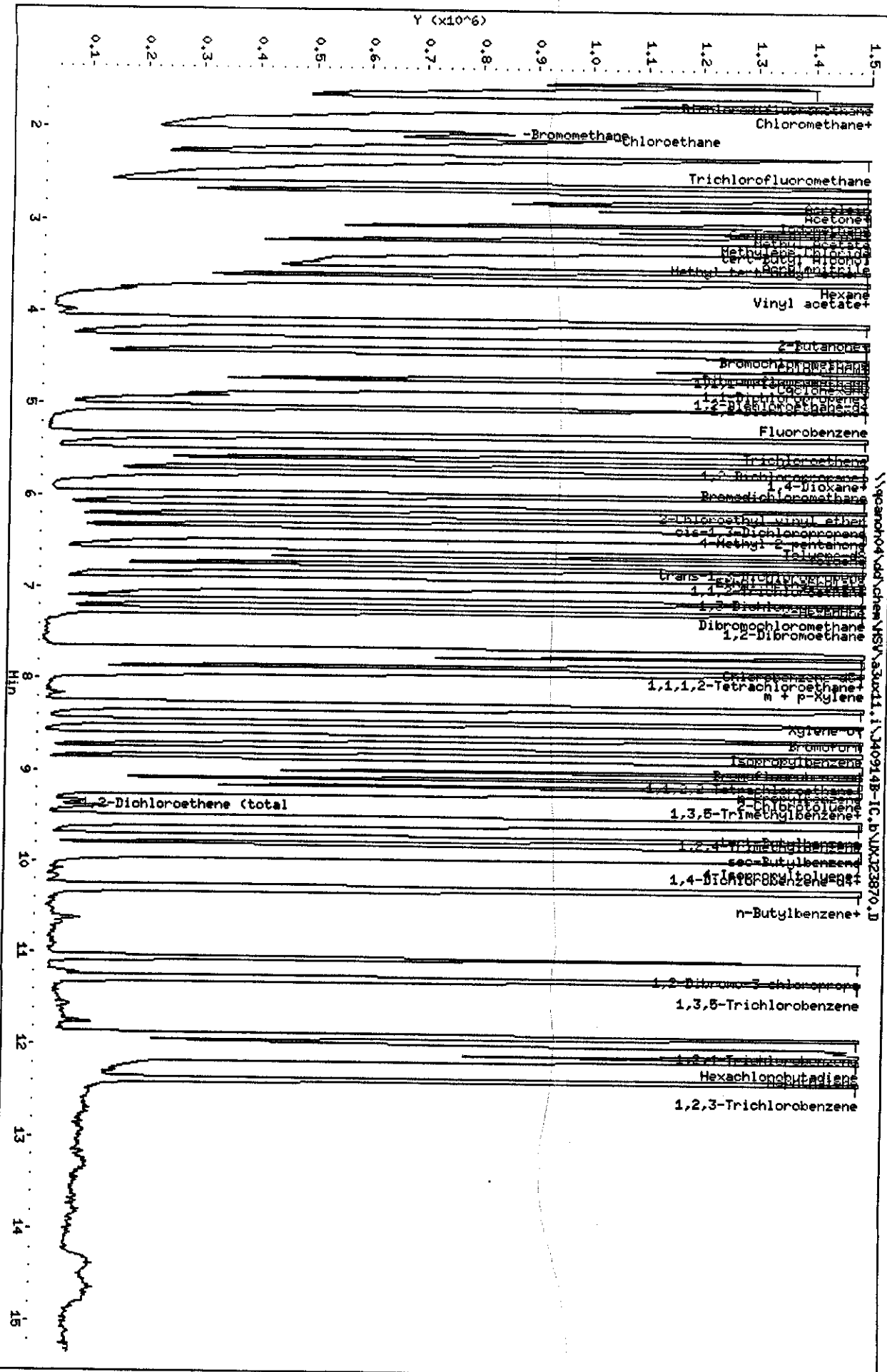
Data File: \\qcanh04\\dd\\chem\\MSV\\a3ux11.i\\40914B-IC.b\\UK323870.D
Date : 14-SEP-2004 13:48
Client ID:
Sample Info: 200NG-IC
Purge Volume: 5.0
Column phase: DB624

Y ($\times 10^6$)

Instrument: a3ux11.i

Operator: 43582

Column diameter: 0.18



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23870.D
Lab Smp Id: 200NG-IC
Inj Date : 14-SEP-2004 13:48
Operator : 43582
Smp Info : 200NG-IC
Misc Info : J40914B-IC,8260LLUX11,2-8260.SUB,43582,1,6
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:45 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 2 Calibration Sample, Level: 6
Dil Factor: 1.00000
Integrator: HP RTE
Target Version: 4.04 Compound Sublist: 2-8260.SUB
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|----------------------------|-----------|---------|-------|---------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * 1 Fluorobenzene | 96 | 5.029 | 5.029 | (1.000) | 2337325 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.668 | 7.668 | (1.000) | 1845289 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 | (1.000) | 1029809 | 50.0000 | | |
| \$ 4 Dibromofluoromethane | 113 | 4.473 | 4.473 | (0.889) | 2180284 | 200.000 | 200.35(A) | |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.745 | 4.745 | (0.944) | 3102729 | 200.000 | 207.35(A) | |
| \$ 6 Toluene-d8 | 98 | 6.366 | 6.366 | (0.830) | 8933577 | 200.000 | 201.52(A) | |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 | (1.145) | 3868072 | 200.000 | 205.36(A) | |
| 8 Dichlorodifluoromethane | 85 | 1.527 | 1.527 | (0.304) | 2468303 | 200.000 | 202.00(A) | |
| 9 Chloromethane | 50 | 1.680 | 1.680 | (0.334) | 4015293 | 200.000 | 184.04 | |
| 10 Vinyl Chloride | 62 | 1.763 | 1.763 | (0.351) | 3096925 | 200.000 | 207.51(A) | |
| 11 Bromomethane | 94 | 2.047 | 2.047 | (0.407) | 1310746 | 200.000 | 185.17 | |
| 12 Chloroethane | 64 | 2.118 | 2.118 | (0.421) | 2147474 | 200.000 | 195.79 | |
| 13 Trichlorofluoromethane | 101 | 2.296 | 2.296 | (0.457) | 3119390 | 200.000 | 199.42 | |
| 15 Acrolein | 56 | 2.603 | 2.603 | (0.518) | 2896520 | 2000.00 | 1983.9 | |
| 16 Acetone | 43 | 2.722 | 2.722 | (0.541) | 1907733 | 400.000 | 400.17(A) | |
| 17 1,1-Dichloroethene | 96 | 2.710 | 2.710 | (0.539) | 2115099 | 200.000 | 202.35(A) | |
| 18 Freon-113 | 151 | 2.722 | 2.722 | (0.541) | 1494012 | 200.000 | 200.47(A) | |

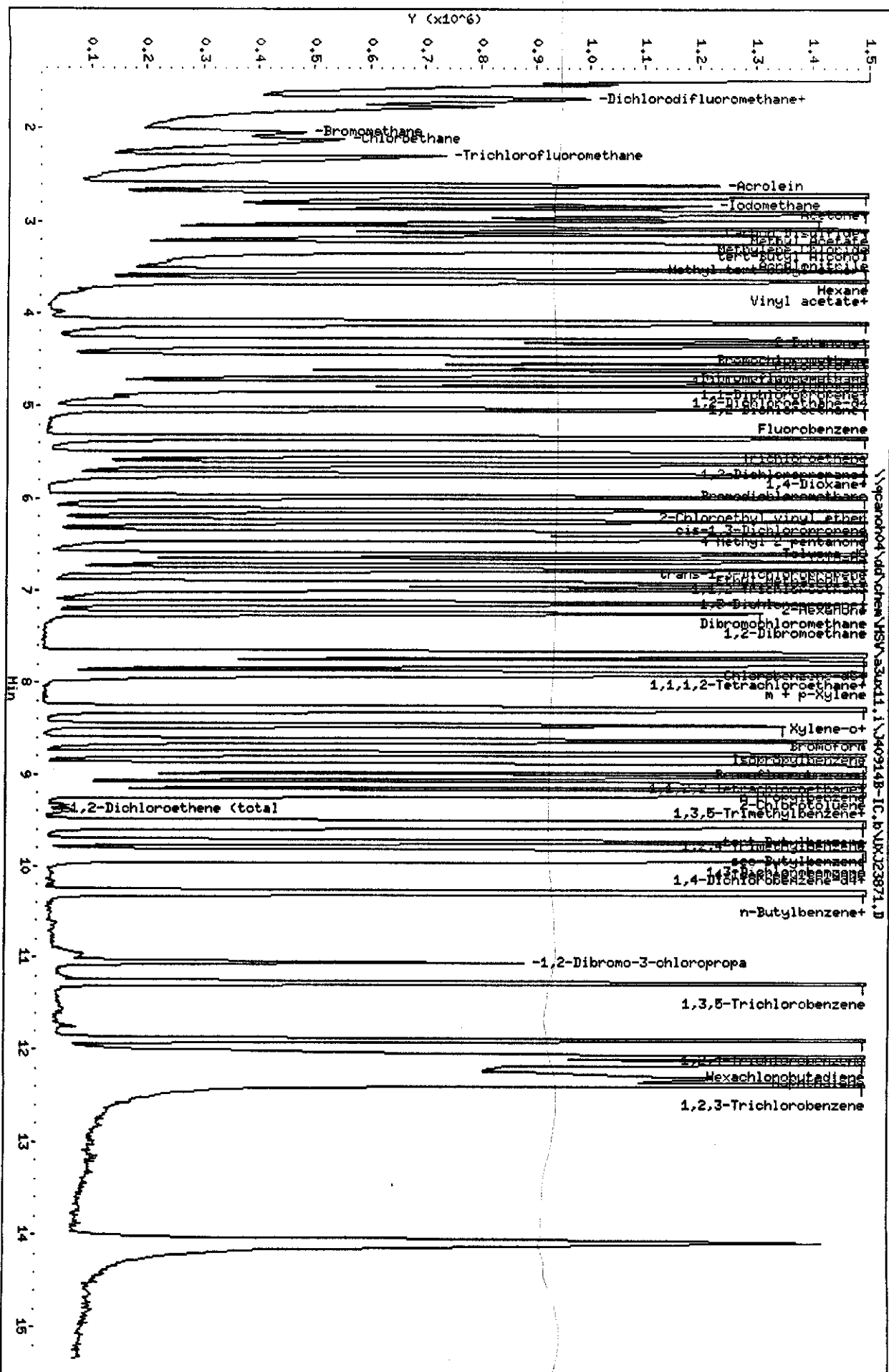
| Compounds | QUANT SIG | | | | | AMOUNTS | | |
|---------------------------------|-----------|-------|--------|---------|----------|------------------|-----------------|--|
| | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) | |
| 19 Iodomethane | 142 | 2.828 | 2.828 | (0.562) | 3111800 | 200.000 | 201.42(A) | |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 | (0.577) | 7890340 | 200.000 | 202.08(A) | |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 | (0.612) | 2622600 | 200.000 | 200.02(A) | |
| 22 Acetonitrile | 41 | 2.935 | 2.935 | (0.584) | 2423653 | 2000.00 | 1747.2 | |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 | (0.647) | 8435321 | 2000.00 | 1955.4 | |
| 24 Methyl tert-butyl ether | 73 | 3.301 | 3.301 | (0.657) | 6861860 | 200.000 | 204.77(A) | |
| 25 trans-1,2-Dichloroethene | 96 | 3.301 | 3.301 | (0.657) | 2403695 | 200.000 | 193.85 | |
| 26 Hexane | 86 | 3.526 | 3.526 | (0.701) | 460062 | 200.000 | 200.35(A) | |
| 27 Vinyl acetate | 43 | 3.656 | 3.656 | (0.727) | 4234296 | 200.000 | 213.40(A) | |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 | (0.722) | 4472069 | 200.000 | 197.70 | |
| 29 tert-Butyl Alcohol | 59 | 3.148 | 3.148 | (0.626) | 3293397 | 4000.00 | 3622.6(A) | |
| 30 2-Butanone | 43 | 4.082 | 4.082 | (0.812) | 2412455 | 400.000 | 376.83(A) | |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 4951179 | 400.000 | 389.38 | |
| 32 cis-1,2-dichloroethene | 96 | 4.094 | 4.094 | (0.814) | 2547484 | 200.000 | 195.53 | |
| 33 2,2-Dichloropropane | 77 | 4.094 | 4.094 | (0.814) | 2760188 | 200.000 | 202.48(A) | |
| 34 Bromochloromethane | 128 | 4.284 | 4.284 | (0.852) | 1158230 | 200.000 | 191.98 | |
| 35 Chloroform | 83 | 4.343 | 4.343 | (0.864) | 4417574 | 200.000 | 194.27 | |
| 36 Tetrahydrofuran | 42 | 4.331 | 4.331 | (0.861) | 658534 | 200.000 | 199.60 | |
| 37 1,1,1-Trichloroethane | 97 | 4.508 | 4.508 | (0.896) | 3459032 | 200.000 | 201.42(A) | |
| 38 1,1-Dichloropropene | 75 | 4.639 | 4.639 | (0.922) | 3312291 | 200.000 | 206.94(A) | |
| 39 Carbon Tetrachloride | 117 | 4.650 | 4.650 | (0.925) | 2829152 | 200.000 | 209.81(A) | |
| 40 1,2-Dichloroethane | 62 | 4.804 | 4.804 | (0.955) | 3659770 | 200.000 | 196.98 | |
| 41 Benzene | 78 | 4.816 | 4.816 | (0.958) | 10307958 | 200.000 | 191.55 | |
| 42 Trichloroethene | 130 | 5.337 | 5.337 | (1.061) | 2454964 | 200.000 | 199.89 | |
| 43 1,2-Dichloropropene | 63 | 5.514 | 5.514 | (1.096) | 2581062 | 200.000 | 194.44 | |
| 44 1,4-Dioxane | 88 | 5.621 | 5.621 | (1.118) | 961094 | 10000.0 | 8446.6(A) | |
| 45 Dibromomethane | 93 | 5.621 | 5.621 | (1.118) | 1453325 | 200.000 | 197.58 | |
| 46 Bromodichloromethane | 83 | 5.739 | 5.739 | (1.141) | 3494837 | 200.000 | 200.14(A) | |
| 47 2-Chloroethyl vinyl ether | 63 | 5.988 | 5.988 | (1.191) | 3196023 | 400.000 | 428.19(A) | |
| 48 cis-1,3-Dichloropropene | 75 | 6.130 | 6.130 | (1.219) | 4425099 | 200.000 | 207.11(A) | |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 | (1.242) | 4708742 | 400.000 | 409.86(A) | |
| 50 Toluene | 91 | 6.425 | 6.425 | (0.838) | 11034583 | 200.000 | 200.23(A) | |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 | (0.861) | 4148070 | 200.000 | 209.93(A) | |
| 52 Ethyl Methacrylate | 69 | 6.674 | 6.674 | (0.870) | 3684757 | 200.000 | 216.74(A) | |
| 53 1,1,2-Trichloroethane | 97 | 6.769 | 6.769 | (0.883) | 2198790 | 200.000 | 196.70 | |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 | (0.903) | 4169551 | 200.000 | 197.96 | |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 | (0.904) | 1776877 | 200.000 | 199.01 | |
| 56 2-Hexanone | 43 | 6.982 | 6.982 | (0.911) | 3736062 | 400.000 | 418.68(A) | |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 | (0.931) | 2430585 | 200.000 | 203.85(A) | |
| 58 1,2-Dibromoethane | 107 | 7.242 | 7.242 | (0.944) | 2251324 | 200.000 | 204.40(A) | |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 | (1.005) | 7055771 | 200.000 | 197.73 | |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 | (1.014) | 2491620 | 200.000 | 200.99(A) | |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 | (1.017) | 3790824 | 200.000 | 210.37(A) | |
| 62 m + p-Xylene | 106 | 7.904 | 7.904 | (1.031) | 9470698 | 400.000 | 408.06(A) | |
| M 63 Xylenes (total) | 106 | | | | 14172021 | 600.000 | 614.39 | |
| 64 Xylene-o | 106 | 8.283 | 8.283 | (1.080) | 4701323 | 200.000 | 206.33(A) | |
| 65 Styrene | 104 | 8.295 | 8.295 | (1.082) | 8616448 | 200.000 | 211.47(A) | |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|--------------------------------|-----------|---------|--------|---------|----------|------------------|-----------------|
| | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| 66 Bromoform | 173 | 8.472 | 8.472 | (1.105) | 1716386 | 200.000 | 211.78 (A) |
| 67 Isopropylbenzene | 105 | 8.626 | 8.626 | (1.125) | 10843610 | 200.000 | 213.68 (A) |
| 68 1,1,2,2-Tetrachloroethane | 83 | 8.898 | 8.898 | (0.898) | 3053504 | 200.000 | 196.70 |
| 69 1,4-Dichloro-2-butene | 53 | 8.958 | 8.958 | (0.904) | 1088280 | 200.000 | 213.07 (A) |
| 70 1,2,3-Trichloropropane | 110 | 8.946 | 8.946 | (0.903) | 994195 | 200.000 | 195.17 |
| 71 Bromobenzene | 156 | 8.922 | 8.922 | (0.901) | 3020894 | 200.000 | 196.61 |
| 72 n-Propylbenzene | 120 | 9.029 | 9.029 | (0.912) | 2967729 | 200.000 | 207.17 (A) |
| 73 2-Chlorotoluene | 126 | 9.111 | 9.111 | (0.920) | 2837950 | 200.000 | 200.05 (A) |
| 74 1,3,5-Trimethylbenzene | 105 | 9.194 | 9.194 | (0.928) | 10066915 | 200.000 | 211.19 (A) |
| 75 4-Chlorotoluene | 126 | 9.218 | 9.218 | (0.931) | 2975348 | 200.000 | 196.33 |
| 76 tert-Butylbenzene | 119 | 9.514 | 9.514 | (0.961) | 7991718 | 200.000 | 206.94 (A) |
| 77 1,2,4-Trimethylbenzene | 105 | 9.561 | 9.561 | (0.965) | 10631151 | 200.000 | 208.92 (A) |
| 78 sec-Butylbenzene | 105 | 9.727 | 9.727 | (0.982) | 10794030 | 200.000 | 206.50 (A) |
| 79 4-Isopropyltoluene | 119 | 9.869 | 9.869 | (0.996) | 9271210 | 200.000 | 213.18 (A) |
| 80 1,3-Dichlorobenzene | 146 | 9.845 | 9.845 | (0.994) | 5493140 | 200.000 | 193.66 |
| 81 1,4-Dichlorobenzene | 146 | 9.928 | 9.928 | (1.002) | 5803120 | 200.000 | 195.54 |
| 82 n-Butylbenzene | 91 | 10.271 | 10.271 | (1.037) | 8332031 | 200.000 | 214.47 (A) |
| 83 1,2-Dichlorobenzene | 146 | 10.295 | 10.295 | (1.039) | 5384330 | 200.000 | 192.72 |
| 84 1,2-Dibromo-3-chloropropane | 157 | 11.052 | 11.052 | (1.116) | 528598 | 200.000 | 201.08 (A) |
| 85 1,2,4-Trichlorobenzene | 180 | 11.892 | 11.892 | (1.201) | 2505741 | 200.000 | 210.67 (A) |
| 86 Hexachlorobutadiene | 225 | 12.070 | 12.070 | (1.219) | 979903 | 200.000 | 200.36 (A) |
| 87 Naphthalene | 128 | 12.129 | 12.129 | (1.225) | 6432622 | 200.000 | 229.14 (A) |
| 88 1,2,3-Trichlorobenzene | 180 | 12.377 | 12.377 | (1.250) | 1787787 | 200.000 | 209.60 (A) |
| 98 Cyclohexane | 56 | 4.568 | 4.568 | (0.908) | 3724145 | 200.000 | 216.58 (A) |
| 143 Methyl Acetate | 43 | 2.982 | 2.982 | (0.593) | 3693009 | 400.000 | 384.10 |
| 144 Methylcyclohexane | 83 | 5.514 | 5.514 | (1.096) | 3054148 | 200.000 | 211.66 |
| 141 1,3,5-Trichlorobenzene | 180 | 11.277 | 11.277 | (1.139) | 3087088 | 200.000 | 196.62 |

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Instrument: 3ux11.i
 Operator: 43582
 Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23871.D
Report Date: 15-Sep-2004 12:45

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23871.D
Lab Smp Id: 100NG-IC
Inj Date : 14-SEP-2004 14:10
Operator : 43582
Smp Info : 100NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 5
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:45 evansl
Cal Date : 14-SEP-2004 15:41
Als bottle: 3
Dil Factor: 1.00000
Integrator: HP RTE
Target Version: 4.04
Processing Host: CANPMSV07
Inst ID: A3UX11.i
Quant Type: ISTD
Cal File: UXJ23875.D
Calibration Sample, Level: 5
Compound Sublist: 2-8260.SUB

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|----------------------------|-----------|---------|-------|---------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * 1 Fluorobenzene | 96 | 5.029 | 5.029 | (1.000) | 2294078 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 | (1.000) | 1821500 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 | (1.000) | 999916 | 50.0000 | | |
| \$ 4 Dibromofluoromethane | 113 | 4.473 | 4.473 | (0.889) | 1074908 | 100.000 | 100.64 | |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.745 | 4.745 | (0.944) | 1555269 | 100.000 | 105.89 | |
| \$ 6 Toluene-d8 | 98 | 6.378 | 6.378 | (0.831) | 4394161 | 100.000 | 100.42 | |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 | (1.143) | 1899831 | 100.000 | 102.18 | |
| 8 Dichlorodifluoromethane | 85 | 1.527 | 1.527 | (0.304) | 1124779 | 100.000 | 93.783 | |
| 9 Chloromethane | 50 | 1.680 | 1.680 | (0.334) | 1934079 | 100.000 | 90.319 | |
| 10 Vinyl Chloride | 62 | 1.763 | 1.763 | (0.351) | 1367463 | 100.000 | 93.356 | |
| 11 Bromomethane | 94 | 2.047 | 2.047 | (0.407) | 634017 | 100.000 | 91.256 | |
| 12 Chloroethane | 64 | 2.130 | 2.130 | (0.424) | 1013498 | 100.000 | 94.147 | |
| 13 Trichlorofluoromethane | 101 | 2.296 | 2.296 | (0.457) | 1404082 | 100.000 | 91.454 | |
| 15 Acrolein | 56 | 2.615 | 2.615 | (0.520) | 1447814 | 1000.00 | 1010.3 | |
| 16 Acetone | 43 | 2.722 | 2.722 | (0.541) | 973885 | 200.000 | 198.82 | |
| 17 1,1-Dichloroethene | 96 | 2.710 | 2.710 | (0.539) | 950397 | 100.000 | 92.638 | |
| 18 Freon-113 | 151 | 2.734 | 2.734 | (0.544) | 619208 | 100.000 | 93.088 | |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|---------------------------------|-----------|---------|-------|---------|---------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| 19 Iodomethane | 142 | 2.828 | 2.828 | (0.562) | 1515000 | 100.000 | 99.909 |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 | (0.577) | 3602731 | 100.000 | 94.011 |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 | (0.612) | 1333682 | 100.000 | 99.833 |
| 22 Acetonitrile | 41 | 2.935 | 2.935 | (0.584) | 1365511 | 1000.00 | 1002.9 |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 | (0.647) | 4234501 | 1000.00 | 1000.1 |
| 24 Methyl tert-butyl ether | 73 | 3.302 | 3.302 | (0.657) | 3440137 | 100.000 | 104.59 |
| 25 trans-1,2-Dichloroethene | 96 | 3.302 | 3.302 | (0.657) | 1175346 | 100.000 | 96.574 |
| 26 Hexane | 86 | 3.526 | 3.526 | (0.701) | 195007 | 100.000 | 95.842 |
| 27 Vinyl acetate | 43 | 3.657 | 3.657 | (0.727) | 2062282 | 100.000 | 105.89 |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 | (0.722) | 2173037 | 100.000 | 97.878 |
| 29 tert-Butyl Alcohol | 59 | 3.148 | 3.148 | (0.626) | 1816490 | 2000.00 | 2035.8(A) |
| 30 2-Butanone | 43 | 4.083 | 4.083 | (0.812) | 1242482 | 200.000 | 197.74 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 2422890 | 200.000 | 194.13 |
| 32 cis-1,2-dichloroethene | 96 | 4.094 | 4.094 | (0.814) | 1247544 | 100.000 | 97.561 |
| 33 2,2-Dichloropropane | 77 | 4.106 | 4.106 | (0.816) | 1285494 | 100.000 | 96.081 |
| 34 Bromochloromethane | 128 | 4.284 | 4.284 | (0.852) | 577210 | 100.000 | 97.480 |
| 35 Chloroform | 83 | 4.343 | 4.343 | (0.864) | 2173231 | 100.000 | 97.374 |
| 36 Tetrahydrofuran | 42 | 4.331 | 4.331 | (0.861) | 344585 | 100.000 | 101.91 |
| 37 1,1,1-Trichloroethane | 97 | 4.509 | 4.509 | (0.896) | 1598802 | 100.000 | 94.853 |
| 38 1,1-Dichloropropene | 75 | 4.639 | 4.639 | (0.922) | 1510060 | 100.000 | 96.123 |
| 39 Carbon Tetrachloride | 117 | 4.651 | 4.651 | (0.925) | 1268602 | 100.000 | 95.854 |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 | (0.958) | 1797145 | 100.000 | 98.552 |
| 41 Benzene | 78 | 4.816 | 4.816 | (0.958) | 5026234 | 100.000 | 95.163 |
| 42 Trichloroethene | 130 | 5.337 | 5.337 | (1.061) | 1185674 | 100.000 | 98.362 |
| 43 1,2-Dichloropropene | 63 | 5.526 | 5.526 | (1.099) | 1270520 | 100.000 | 97.515 |
| 44 1,4-Dioxane | 88 | 5.621 | 5.621 | (1.118) | 597256 | 5000.00 | 5347.9(A) |
| 45 Dibromomethane | 93 | 5.621 | 5.621 | (1.118) | 727774 | 100.000 | 100.80 |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 | (1.144) | 1694447 | 100.000 | 98.863 |
| 47 2-Chloroethyl vinyl ether | 63 | 5.988 | 5.988 | (1.191) | 1586020 | 200.000 | 216.50(A) |
| 48 cis-1,3-Dichloropropene | 75 | 6.130 | 6.130 | (1.219) | 2139621 | 100.000 | 102.03 |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 | (1.242) | 2343412 | 200.000 | 207.82(A) |
| 50 Toluene | 91 | 6.437 | 6.437 | (0.838) | 5341010 | 100.000 | 98.181 |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 | (0.860) | 2025722 | 100.000 | 103.86 |
| 52 Ethyl Methacrylate | 69 | 6.674 | 6.674 | (0.869) | 1801585 | 100.000 | 107.35 |
| 53 1,1,2-Trichloroethane | 97 | 6.769 | 6.769 | (0.881) | 1103061 | 100.000 | 99.965 |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 | (0.901) | 2083135 | 100.000 | 100.19 |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 | (0.903) | 827625 | 100.000 | 93.905 |
| 56 2-Hexanone | 43 | 6.982 | 6.982 | (0.909) | 1792800 | 200.000 | 203.53(A) |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 | (0.929) | 1188499 | 100.000 | 100.98 |
| 58 1,2-Dibromoethane | 107 | 7.242 | 7.242 | (0.943) | 1119552 | 100.000 | 102.98 |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 | (1.003) | 3458216 | 100.000 | 98.180 |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 | (1.012) | 1239765 | 100.000 | 101.31 |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 | (1.015) | 1782706 | 100.000 | 100.22 |
| 62 m + p-Xylene | 106 | 7.905 | 7.905 | (1.029) | 4519264 | 200.000 | 197.26 |
| M 63 Xylenes (total) | 106 | | | | 6782922 | 300.000 | 297.91 |
| 64 Xylene-o | 106 | 8.283 | 8.283 | (1.079) | 2263658 | 100.000 | 100.65 |
| 65 Styrene | 104 | 8.295 | 8.295 | (1.080) | 4147114 | 100.000 | 103.11 |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|--------------------------------|-----------|---------|--------|---------|---------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| 66 Bromoform | 173 | 8.473 | 8.473 | (1.103) | 820790 | 100.000 | 102.60 |
| 67 Isopropylbenzene | 105 | 8.626 | 8.626 | (1.123) | 4943240 | 100.000 | 98.682 |
| 68 1,1,2,2-Tetrachloroethane | 83 | 8.899 | 8.899 | (0.898) | 1522185 | 100.000 | 100.99 |
| 69 1,4-Dichloro-2-butene | 53 | 8.958 | 8.958 | (0.904) | 523620 | 100.000 | 105.58 |
| 70 1,2,3-Trichloropropane | 110 | 8.946 | 8.946 | (0.903) | 495410 | 100.000 | 100.16 |
| 71 Bromobenzene | 156 | 8.934 | 8.934 | (0.902) | 1490126 | 100.000 | 99.884 |
| 72 n-Propylbenzene | 120 | 9.029 | 9.029 | (0.912) | 1351668 | 100.000 | 97.179 |
| 73 2-Chlorotoluene | 126 | 9.112 | 9.112 | (0.920) | 1351055 | 100.000 | 98.084 |
| 74 1,3,5-Trimethylbenzene | 105 | 9.194 | 9.194 | (0.928) | 4630795 | 100.000 | 100.05 |
| 75 4-Chlorotoluene | 126 | 9.218 | 9.218 | (0.931) | 1433771 | 100.000 | 97.438 |
| 76 tert-Butylbenzene | 119 | 9.514 | 9.514 | (0.961) | 3670015 | 100.000 | 97.876 |
| 77 1,2,4-Trimethylbenzene | 105 | 9.561 | 9.561 | (0.965) | 5001495 | 100.000 | 101.23 |
| 78 sec-Butylbenzene | 105 | 9.727 | 9.727 | (0.982) | 4894571 | 100.000 | 96.437 |
| 79 4-Isopropyltoluene | 119 | 9.869 | 9.869 | (0.996) | 4211412 | 100.000 | 99.734 |
| 80 1,3-Dichlorobenzene | 146 | 9.845 | 9.845 | (0.994) | 2672507 | 100.000 | 97.033 |
| 81 1,4-Dichlorobenzene | 146 | 9.928 | 9.928 | (1.002) | 2817287 | 100.000 | 97.770 |
| 82 n-Butylbenzene | 91 | 10.271 | 10.271 | (1.037) | 3687678 | 100.000 | 97.759 |
| 83 1,2-Dichlorobenzene | 146 | 10.295 | 10.295 | (1.039) | 2647056 | 100.000 | 97.579 |
| 84 1,2-Dibromo-3-chloropropane | 157 | 11.052 | 11.052 | (1.116) | 261375 | 100.000 | 102.40 |
| 85 1,2,4-Trichlorobenzene | 180 | 11.892 | 11.892 | (1.201) | 1158323 | 100.000 | 100.30 |
| 86 Hexachlorobutadiene | 225 | 12.070 | 12.070 | (1.219) | 475139 | 100.000 | 97.147 |
| 87 Naphthalene | 128 | 12.129 | 12.129 | (1.225) | 3076588 | 100.000 | 112.87 |
| 88 1,2,3-Trichlorobenzene | 180 | 12.377 | 12.377 | (1.250) | 883388 | 100.000 | 106.66 |
| 98 Cyclohexane | 56 | 4.568 | 4.568 | (0.908) | 1591353 | 100.000 | 94.291 |
| 143 Methyl Acetate | 43 | 2.994 | 2.994 | (0.595) | 1827750 | 200.000 | 193.68 |
| 144 Methylcyclohexane | 83 | 5.514 | 5.514 | (1.096) | 1298110 | 100.000 | 91.659 |
| 141 1,3,5-Trichlorobenzene | 180 | 11.277 | 11.277 | (1.139) | 1469129 | 100.000 | 96.368 |

QC Flag Legend

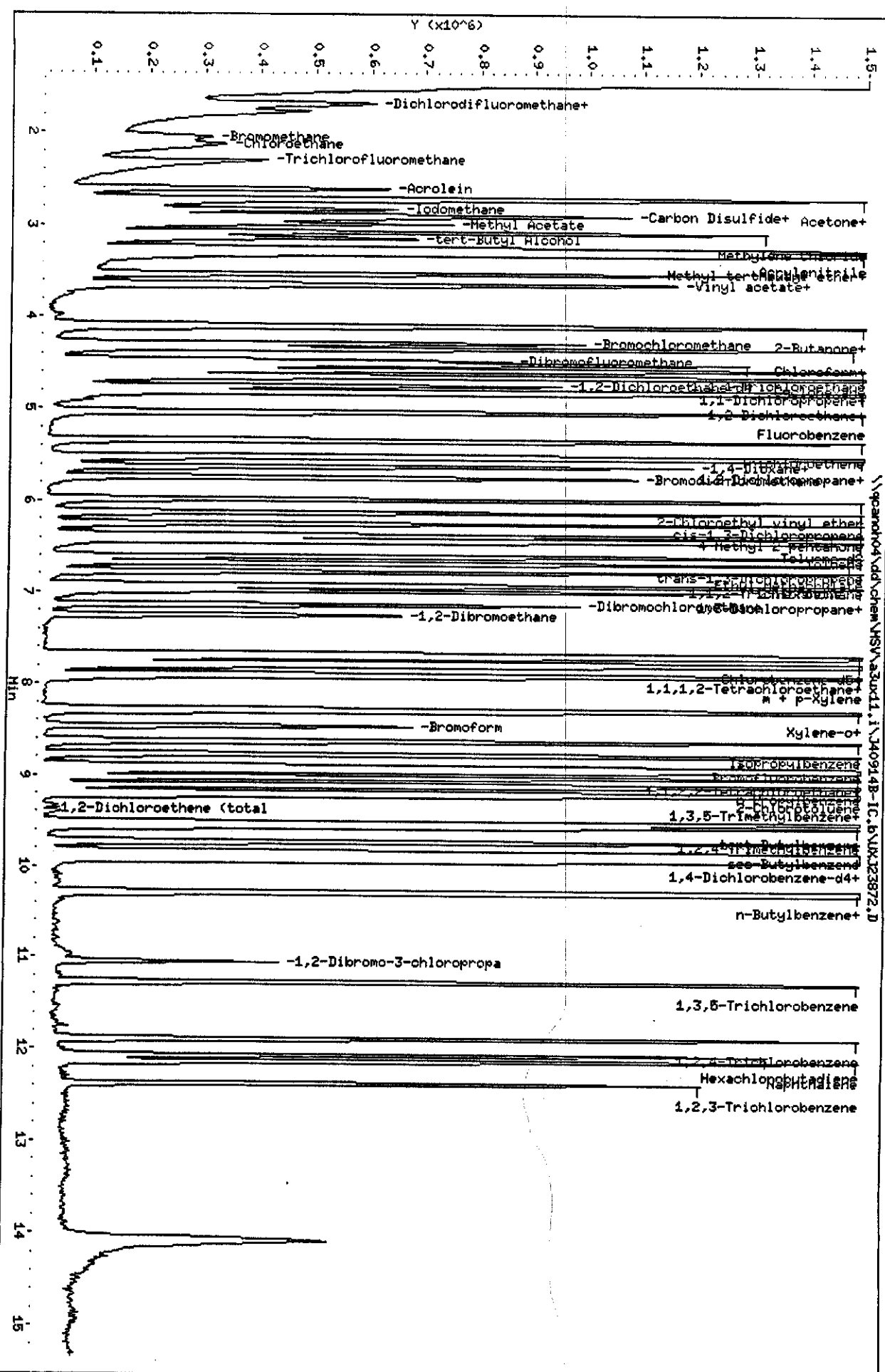
A - Target compound detected but, quantitated amount exceeded maximum amount.

Client ID:
 Sample Info: 50mg-IC
 Purge Volume: 5.0

Column phase: DB624

Instrument: 30x11.i

Operator: 43582
 Column diameter: 0.18



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23872.D
Lab Smp Id: 50NG-IC
Inj Date : 14-SEP-2004 14:33
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 50NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 4
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:46 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 4 Calibration Sample, Level: 4
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|-----------|--------------------------|---------|-------|---------------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * | 1 Fluorobenzene | 96 | 5.029 | 5.029 (1.000) | 2329625 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.668 | 7.668 (1.000) | 1858993 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 990958 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.473 | 4.473 (0.889) | 558177 | 50.0000 | | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.745 | 4.745 (0.944) | 733545 | 50.0000 | 49.183 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.832) | 2256469 | 50.0000 | 50.525 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.145) | 943942 | 50.0000 | 49.744 | |
| 8 | Dichlorodifluoromethane | 85 | 1.527 | 1.527 (0.304) | 586151 | 50.0000 | 48.127 | |
| 9 | Chloromethane | 50 | 1.680 | 1.680 (0.334) | 1022123 | 50.0000 | 47.004 | |
| 10 | Vinyl Chloride | 62 | 1.763 | 1.763 (0.351) | 726372 | 50.0000 | 48.832 | |
| 11 | Bromomethane | 94 | 2.047 | 2.047 (0.407) | 352300 | 50.0000 | 49.934 | |
| 12 | Chloroethane | 64 | 2.130 | 2.130 (0.424) | 545580 | 50.0000 | 49.907 | |
| 13 | Trichlorofluoromethane | 101 | 2.296 | 2.296 (0.457) | 757893 | 50.0000 | 48.612 | |
| 15 | Acrolein | 56 | 2.615 | 2.615 (0.520) | 720539 | 500.000 | 495.15 | |
| 16 | Acetone | 43 | 2.722 | 2.722 (0.541) | 530389 | 100.000 | 102.36 | |
| 17 | 1,1-Dichloroethene | 96 | 2.710 | 2.710 (0.539) | 541982 | 50.0000 | 52.022 | |
| 18 | Freon-113 | 151 | 2.722 | 2.722 (0.541) | 390502 | 50.0000 | 59.593 | |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|---------------------------------|-----------|---------|-------|---------|---------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| 19 Iodomethane | 142 | 2.828 | 2.828 | (0.562) | 777257 | 50.0000 | 50.475 |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 | (0.577) | 1989456 | 50.0000 | 51.121 |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 | (0.612) | 741159 | 50.0000 | 50.431 |
| 22 Acetonitrile | 41 | 2.935 | 2.935 | (0.584) | 667504 | 500.000 | 482.78 |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 | (0.647) | 2117146 | 500.000 | 492.41 |
| 24 Methyl tert-butyl ether | 73 | 3.302 | 3.302 | (0.657) | 1733984 | 50.0000 | 51.915 |
| 25 trans-1,2-Dichloroethene | 96 | 3.302 | 3.302 | (0.657) | 623149 | 50.0000 | 50.421 |
| 26 Hexane | 86 | 3.526 | 3.526 | (0.701) | 115096 | 50.0000 | 57.272 |
| 27 Vinyl acetate | 43 | 3.657 | 3.657 | (0.727) | 979072 | 50.0000 | 49.506 |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 | (0.722) | 1126418 | 50.0000 | 49.962 |
| 29 tert-Butyl Alcohol | 59 | 3.148 | 3.148 | (0.626) | 896352 | 1000.00 | 989.22 |
| 30 2-Butanone | 43 | 4.083 | 4.083 | (0.812) | 623229 | 100.000 | 97.672 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 1262694 | 100.000 | 99.671 |
| 32 cis-1,2-dichloroethene | 96 | 4.094 | 4.094 | (0.814) | 639545 | 50.0000 | 49.251 |
| 33 2,2-Dichloropropane | 77 | 4.106 | 4.106 | (0.816) | 684525 | 50.0000 | 50.382 |
| 34 Bromochloromethane | 128 | 4.284 | 4.284 | (0.852) | 299703 | 50.0000 | 49.842 |
| 35 Chloroform | 83 | 4.343 | 4.343 | (0.864) | 1133371 | 50.0000 | 50.007 |
| 36 Tetrahydrofuran | 42 | 4.331 | 4.331 | (0.861) | 170087 | 50.0000 | 48.338 |
| 37 1,1,1-Trichloroethane | 97 | 4.508 | 4.508 | (0.896) | 892252 | 50.0000 | 52.127 |
| 38 1,1-Dichloropropene | 75 | 4.639 | 4.639 | (0.922) | 833818 | 50.0000 | 52.267 |
| 39 Carbon Tetrachloride | 117 | 4.650 | 4.650 | (0.925) | 724043 | 50.0000 | 53.873 |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 | (0.958) | 914079 | 50.0000 | 49.362 |
| 41 Benzene | 78 | 4.816 | 4.816 | (0.958) | 2642144 | 50.0000 | 49.261 |
| 42 Trichloroethene | 130 | 5.337 | 5.337 | (1.061) | 622755 | 50.0000 | 50.875 |
| 43 1,2-Dichloropropene | 63 | 5.526 | 5.526 | (1.099) | 654580 | 50.0000 | 49.474 |
| 44 1,4-Dioxane | 88 | 5.621 | 5.621 | (1.118) | 304150 | 2500.00 | 2681.8 (A) |
| 45 Dibromomethane | 93 | 5.621 | 5.621 | (1.118) | 368591 | 50.0000 | 50.275 |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 | (1.144) | 862931 | 50.0000 | 49.580 |
| 47 2-Chloroethyl vinyl ether | 63 | 5.988 | 5.988 | (1.191) | 777773 | 100.000 | 104.55 |
| 48 cis-1,3-Dichloropropene | 75 | 6.130 | 6.130 | (1.219) | 1059681 | 50.0000 | 49.761 |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 | (1.242) | 1174753 | 100.000 | 102.59 |
| 50 Toluene | 91 | 6.437 | 6.437 | (0.840) | 2773477 | 50.0000 | 49.955 |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 | (0.861) | 1008249 | 50.0000 | 50.651 |
| 52 Ethyl Methacrylate | 69 | 6.674 | 6.674 | (0.870) | 893443 | 50.0000 | 52.165 |
| 53 1,1,2-Trichloroethane | 97 | 6.769 | 6.769 | (0.883) | 564234 | 50.0000 | 50.102 |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 | (0.903) | 1040578 | 50.0000 | 49.040 |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 | (0.904) | 455621 | 50.0000 | 50.654 |
| 56 2-Hexanone | 43 | 6.982 | 6.982 | (0.911) | 925357 | 100.000 | 102.93 |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 | (0.931) | 593974 | 50.0000 | 49.450 |
| 58 1,2-Dibromoethane | 107 | 7.242 | 7.242 | (0.944) | 561972 | 50.0000 | 50.647 |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 | (1.005) | 1760577 | 50.0000 | 48.975 |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 | (1.014) | 612067 | 50.0000 | 49.009 |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 | (1.017) | 909162 | 50.0000 | 50.081 |
| 62 m + p-Xylene | 106 | 7.905 | 7.905 | (1.031) | 2376751 | 100.000 | 101.65 |
| M 63 Xylenes (total) | 106 | 8.283 | 8.283 | (1.080) | 3528868 | 150.000 | 151.84 |
| 64 Xylene-o | 106 | 8.295 | 8.295 | (1.082) | 1152117 | 50.0000 | 50.192 |
| 65 Styrene | 104 | 8.295 | 8.295 | (1.082) | 2072070 | 50.0000 | 50.479 |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|--------------------------------|-----------|---------|--------|---------|--------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| 66 Bromoform | 173 | 8.473 | 8.473 | (1.105) | | 406002 | 50.0000 | 49.727 |
| 67 Isopropylbenzene | 105 | 8.626 | 8.626 | (1.125) | | 2624726 | 50.0000 | 51.341 |
| 68 1,1,2,2-Tetrachloroethane | 83 | 8.899 | 8.899 | (0.898) | | 750188 | 50.0000 | 50.220 |
| 69 1,4-Dichloro-2-butene | 53 | 8.958 | 8.958 | (0.904) | | 249440 | 50.0000 | 50.752 |
| 70 1,2,3-Trichloropropane | 110 | 8.946 | 8.946 | (0.903) | | 246053 | 50.0000 | 50.196 |
| 71 Bromobenzene | 156 | 8.934 | 8.934 | (0.902) | | 745059 | 50.0000 | 50.393 |
| 72 n-Propylbenzene | 120 | 9.029 | 9.029 | (0.912) | | 716364 | 50.0000 | 51.969 |
| 73 2-Chlorotoluene | 126 | 9.111 | 9.111 | (0.920) | | 694404 | 50.0000 | 50.868 |
| 74 1,3,5-Trimethylbenzene | 105 | 9.194 | 9.194 | (0.928) | | 2372465 | 50.0000 | 51.722 |
| 75 4-Chlorotoluene | 126 | 9.218 | 9.218 | (0.931) | | 738082 | 50.0000 | 50.613 |
| 76 tert-Butylbenzene | 119 | 9.514 | 9.514 | (0.961) | | 1923678 | 50.0000 | 51.766 |
| 77 1,2,4-Trimethylbenzene | 105 | 9.561 | 9.561 | (0.965) | | 2551790 | 50.0000 | 52.114 |
| 78 sec-Butylbenzene | 105 | 9.727 | 9.727 | (0.982) | | 2633655 | 50.0000 | 52.360 |
| 79 4-Isopropyltoluene | 119 | 9.869 | 9.869 | (0.996) | | 2250860 | 50.0000 | 53.786 |
| 80 1,3-Dichlorobenzene | 146 | 9.845 | 9.845 | (0.994) | | 1354905 | 50.0000 | 49.638 |
| 81 1,4-Dichlorobenzene | 146 | 9.928 | 9.928 | (1.002) | | 1437001 | 50.0000 | 50.320 |
| 82 n-Butylbenzene | 91 | 10.271 | 10.271 | (1.037) | | 2001622 | 50.0000 | 53.542 |
| 83 1,2-Dichlorobenzene | 146 | 10.295 | 10.295 | (1.039) | | 1339377 | 50.0000 | 49.820 |
| 84 1,2-Dibromo-3-chloropropane | 157 | 11.052 | 11.052 | (1.116) | | 127154 | 50.0000 | 50.267 |
| 85 1,2,4-Trichlorobenzene | 180 | 11.892 | 11.892 | (1.201) | | 576092 | 50.0000 | 50.333 |
| 86 Hexachlorobutadiene | 225 | 12.070 | 12.070 | (1.219) | | 272701 | 50.0000 | 54.568 |
| 87 Naphthalene | 128 | 12.129 | 12.129 | (1.225) | | 1382075 | 50.0000 | 51.162 |
| 88 1,2,3-Trichlorobenzene | 180 | 12.377 | 12.377 | (1.250) | | 415515 | 50.0000 | 50.625 |
| 98 Cyclohexane | 56 | 4.568 | 4.568 | (0.908) | | 952028 | 50.0000 | 55.549 |
| 143 Methyl Acetate | 43 | 2.994 | 2.994 | (0.595) | | 911479 | 100.000 | 95.114 |
| 144 Methylcyclohexane | 83 | 5.514 | 5.514 | (1.096) | | 793891 | 50.0000 | 55.200 |
| 141 1,3,5-Trichlorobenzene | 180 | 11.277 | 11.277 | (1.139) | | 752113 | 50.0000 | 49.781 |

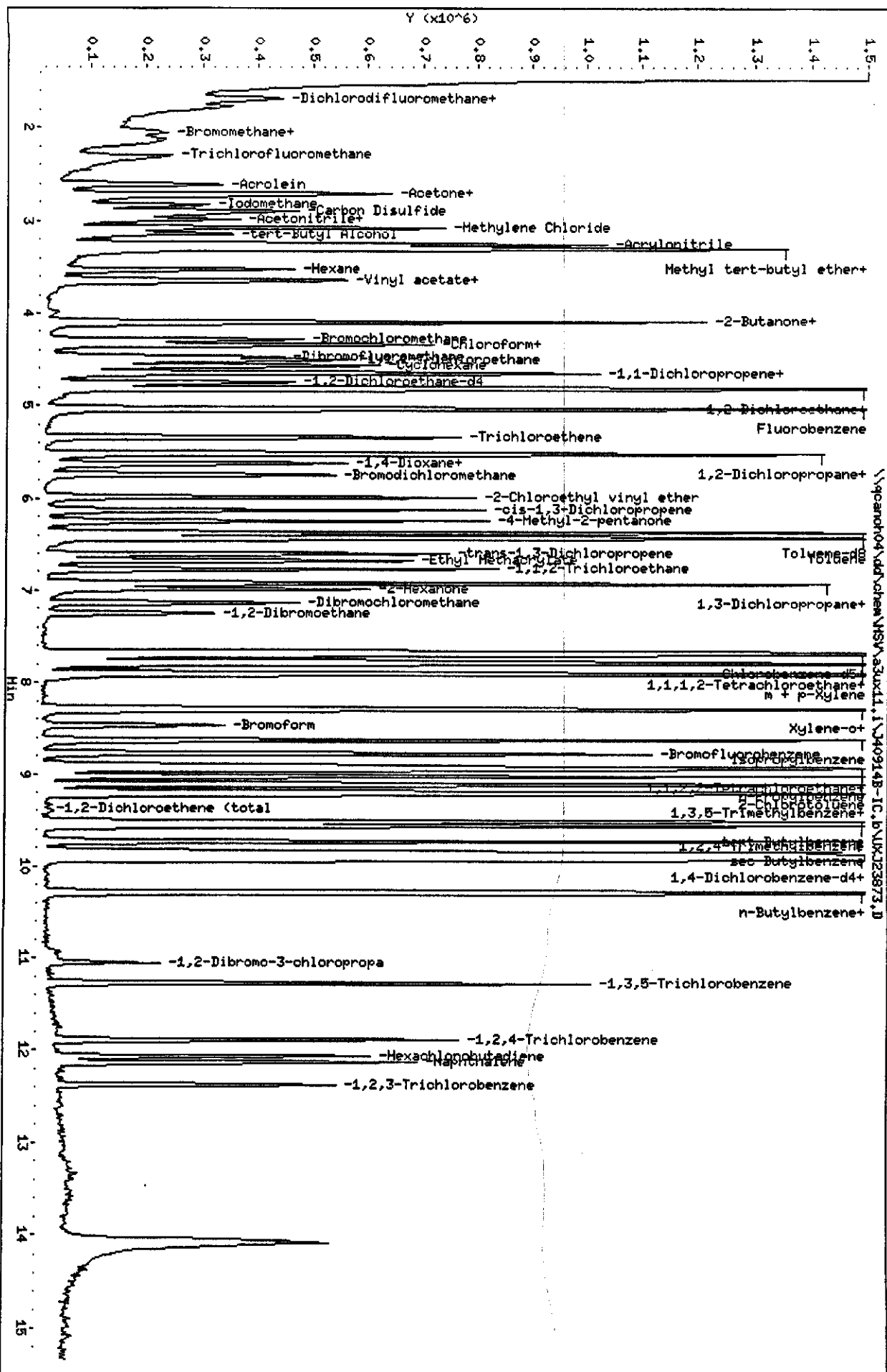
QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Instrument: a3ux11.i

Operator: 43582

Column diameter: 0.18
 \\qcach04\\dd\\ches\\MSV\\a3ux11.i\\40914B-IC.b\\UKJ23873.D



STL North Canton

VOLATILE REPORT SW-846 Method
Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23873.D
Lab Smp Id: 25NG-IC
Inj Date : 14-SEP-2004 14:57
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 25NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 3
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:46 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 5 Calibration Sample, Level: 3
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|----------------------------|-----------|---------|---------------|--------|---------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| * 1 Fluorobenzene | 96 | 5.029 | 5.029 (1.000) | | 2302064 | 50.0000 | |
| * 2 Chlorobenzene-d5 | 117 | 7.668 | 7.668 (1.000) | | 1820780 | 50.0000 | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | | 1002006 | 50.0000 | |
| \$ 4 Dibromofluoromethane | 113 | 4.473 | 4.473 (0.889) | | 263579 | 25.0000 | 24.592 |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.745 | 4.745 (0.944) | | 360514 | 25.0000 | 24.461 |
| \$ 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.832) | | 1137706 | 25.0000 | 26.009 |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.145) | | 468289 | 25.0000 | 25.196 |
| 8 Dichlorodifluoromethane | 85 | 1.526 | 1.526 (0.304) | | 324283 | 25.0000 | 26.945 |
| 9 Chloromethane | 50 | 1.680 | 1.680 (0.334) | | 523520 | 25.0000 | 24.363 |
| 10 Vinyl Chloride | 62 | 1.775 | 1.775 (0.353) | | 382674 | 25.0000 | 26.034 |
| 11 Bromomethane | 94 | 2.047 | 2.047 (0.407) | | 163334 | 25.0000 | 23.428 |
| 12 Chloroethane | 64 | 2.130 | 2.130 (0.424) | | 280413 | 25.0000 | 25.958 |
| 13 Trichlorofluoromethane | 101 | 2.296 | 2.296 (0.457) | | 399606 | 25.0000 | 25.938 |
| 15 Acrolein | 56 | 2.615 | 2.615 (0.520) | | 360127 | 250.000 | 250.44 |
| 16 Acetone | 43 | 2.722 | 2.722 (0.541) | | 262942 | 50.0000 | 48.200 |
| 17 1,1-Dichloroethene | 96 | 2.710 | 2.710 (0.539) | | 229496 | 25.0000 | 22.292 |
| 18 Freon-113 | 151 | 2.722 | 2.722 (0.541) | | 126246 | 25.0000 | 19.380 |

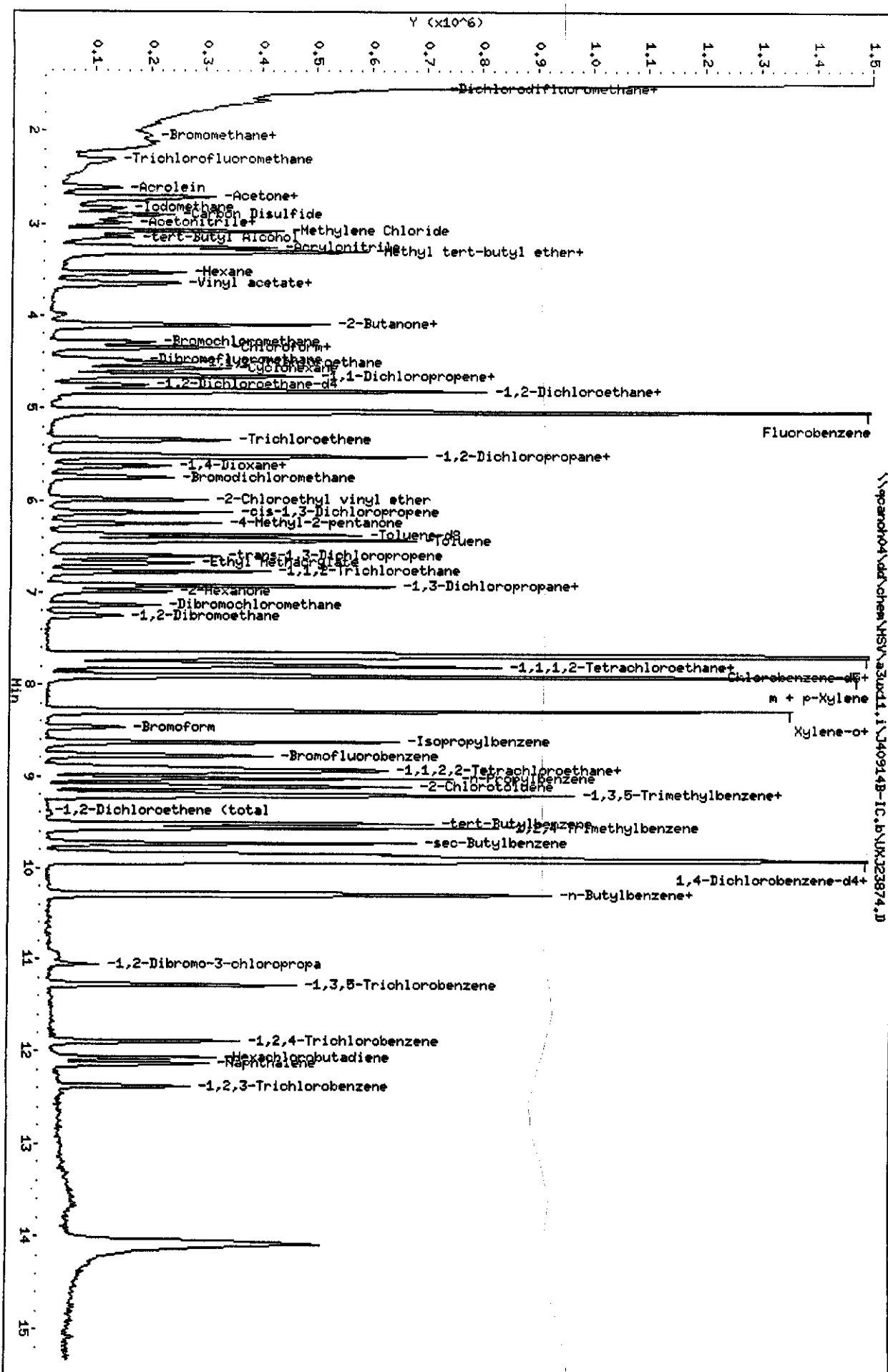
| Compounds | QUANT SIG | | | | | AMOUNTS | |
|---------------------------------|-----------|-------|---------------|--------|----------|------------------|-----------------|
| | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| 19 Iodomethane | 142 | 2.828 | 2.828 (0.562) | | 375596 | 25.0000 | 24.683 |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 (0.577) | | 854870 | 25.0000 | 22.230 |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 (0.612) | | 413458 | 25.0000 | 24.263 |
| 22 Acetonitrile | 41 | 2.946 | 2.946 (0.586) | | 352454 | 250.000 | 257.97 |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 (0.647) | | 1072529 | 250.000 | 252.44 |
| 24 Methyl tert-butyl ether | 73 | 3.301 | 3.301 (0.657) | | 855584 | 25.0000 | 25.923 |
| 25 trans-1,2-Dichloroethene | 96 | 3.313 | 3.313 (0.659) | | 291530 | 25.0000 | 23.871 |
| 26 Hexane | 86 | 3.526 | 3.526 (0.701) | | 38256 | 25.0000 | 19.484 |
| 27 Vinyl acetate | 43 | 3.656 | 3.656 (0.727) | | 490495 | 25.0000 | 25.099 |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 (0.722) | | 530330 | 25.0000 | 23.804 |
| 29 tert-Butyl Alcohol | 59 | 3.148 | 3.148 (0.626) | | 445116 | 500.000 | 497.11 |
| 30 2-Butanone | 43 | 4.082 | 4.082 (0.812) | | 314848 | 50.0000 | 49.934 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 603009 | 50.0000 | 48.145 |
| 32 cis-1,2-dichloroethene | 96 | 4.094 | 4.094 (0.814) | | 311479 | 25.0000 | 24.274 |
| 33 2,2-Dichloropropane | 77 | 4.106 | 4.106 (0.816) | | 318520 | 25.0000 | 23.724 |
| 34 Bromochloromethane | 128 | 4.284 | 4.284 (0.852) | | 145392 | 25.0000 | 24.469 |
| 35 Chloroform | 83 | 4.343 | 4.343 (0.864) | | 541044 | 25.0000 | 24.158 |
| 36 Tetrahydrofuran | 42 | 4.331 | 4.331 (0.861) | | 78777 | 25.0000 | 22.465 |
| 37 1,1,1-Trichloroethane | 97 | 4.508 | 4.508 (0.896) | | 386062 | 25.0000 | 22.825 |
| 38 1,1-Dichloropropene | 75 | 4.639 | 4.639 (0.922) | | 352859 | 25.0000 | 22.383 |
| 39 Carbon Tetrachloride | 117 | 4.650 | 4.650 (0.925) | | 282428 | 25.0000 | 21.266 |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 (0.958) | | 443026 | 25.0000 | 24.210 |
| 41 Benzene | 78 | 4.816 | 4.816 (0.958) | | 1294581 | 25.0000 | 24.426 |
| 42 Trichloroethene | 130 | 5.337 | 5.337 (1.061) | | 289238 | 25.0000 | 23.912 |
| 43 1,2-Dichloropropane | 63 | 5.526 | 5.526 (1.099) | | 313009 | 25.0000 | 23.941 |
| 44 1,4-Dioxane | 88 | 5.621 | 5.621 (1.118) | | 148470 | 1250.00 | 1324.8 (A) |
| 45 Dibromomethane | 93 | 5.621 | 5.621 (1.118) | | 175336 | 25.0000 | 24.202 |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 (1.144) | | 415519 | 25.0000 | 24.160 |
| 47 2-Chloroethyl vinyl ether | 63 | 5.987 | 5.987 (1.191) | | 367975 | 50.0000 | 50.055 |
| 48 cis-1,3-Dichloropropene | 75 | 6.129 | 6.129 (1.219) | | 502234 | 25.0000 | 23.866 |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 (1.242) | | 557786 | 50.0000 | 49.294 |
| 50 Toluene | 91 | 6.437 | 6.437 (0.840) | | 1338715 | 25.0000 | 24.618 |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 (0.861) | | 468202 | 25.0000 | 24.015 |
| 52 Ethyl Methacrylate | 69 | 6.674 | 6.674 (0.870) | | 409677 | 25.0000 | 24.422 |
| 53 1,1,2-Trichloroethane | 97 | 6.768 | 6.768 (0.883) | | 270822 | 25.0000 | 24.553 |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 (0.903) | | 513387 | 25.0000 | 24.702 |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 (0.904) | | | | |
| 56 2-Hexanone | 43 | 6.981 | 6.981 (0.911) | | 201576 | 25.0000 | 22.880 |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 (0.931) | | 441195 | 50.0000 | 50.107 |
| 58 1,2-Dibromoethane | 107 | 7.254 | 7.254 (0.946) | | 291299 | 25.0000 | 24.760 |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 (1.005) | | 264386 | 25.0000 | 24.328 |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 (1.014) | | 859376 | 25.0000 | 24.408 |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 (1.017) | | 297129 | 25.0000 | 24.291 |
| 62 m + p-Xylene | 106 | 7.904 | 7.904 (1.031) | | 1125922 | 50.0000 | 49.164 |
| M 63 Xylenes (total) | 106 | | | | 1692886 | 75.0000 | 74.383 |
| 64 Xylene-o | 106 | 8.283 | 8.283 (1.080) | | 566964 | 25.0000 | 25.218 |
| 65 Styrene | 104 | 8.295 | 8.295 (1.082) | | 985947 | 25.0000 | 24.523 |

| Compounds | QUANT SIG | MASS | RT | EXP RT | REL RT | RESPONSE | AMOUNTS | |
|--------------------------------|-----------|------|--------|----------------|--------|----------|------------------|-----------------|
| | | | | | | | CAL-AMT (ng) | ON-COL (ng) |
| 66 Bromoform | | 173 | 8.472 | 8.472 (1.105) | | 195832 | 25.0000 | 24.489 |
| 67 Isopropylbenzene | | 105 | 8.626 | 8.626 (1.125) | | 1187991 | 25.0000 | 23.725 |
| 68 1,1,2,2-Tetrachloroethane | | 83 | 8.898 | 8.898 (0.898) | | 370501. | 25.0000 | 24.529 |
| 69 1,4-Dichloro-2-butene | | 53 | 8.958 | 8.958 (0.904) | | 119797 | 25.0000 | 24.105 |
| 70 1,2,3-Trichloropropane | | 110 | 8.946 | 8.946 (0.903) | | 121026 | 25.0000 | 24.417 |
| 71 Bromobenzene | | 156 | 8.922 | 8.922 (0.901) | | 361498 | 25.0000 | 24.181 |
| 72 n-Propylbenzene | | 120 | 9.029 | 9.029 (0.912) | | 315155 | 25.0000 | 22.611 |
| 73 2-Chlorotoluene | | 126 | 9.111 | 9.111 (0.920) | | 330096 | 25.0000 | 23.914 |
| 74 1,3,5-Trimethylbenzene | | 105 | 9.194 | 9.194 (0.928) | | 1104880 | 25.0000 | 23.822 |
| 75 4-Chlorotoluene | | 126 | 9.218 | 9.218 (0.931) | | 353698 | 25.0000 | 23.987 |
| 76 tert-Butylbenzene | | 119 | 9.514 | 9.514 (0.961) | | 852078 | 25.0000 | 22.677 |
| 77 1,2,4-Trimethylbenzene | | 105 | 9.561 | 9.561 (0.965) | | 1171362 | 25.0000 | 23.658 |
| 78 sec-Butylbenzene | | 105 | 9.727 | 9.727 (0.982) | | 1145024 | 25.0000 | 22.513 |
| 79 4-Isopropyltoluene | | 119 | 9.869 | 9.869 (0.996) | | 954488 | 25.0000 | 22.557 |
| 80 1,3-Dichlorobenzene | | 146 | 9.845 | 9.845 (0.994) | | 654290 | 25.0000 | 23.706 |
| 81 1,4-Dichlorobenzene | | 146 | 9.928 | 9.928 (1.002) | | 690846 | 25.0000 | 23.925 |
| 82 n-Butylbenzene | | 91 | 10.271 | 10.271 (1.037) | | 825372 | 25.0000 | 21.835 |
| 83 1,2-Dichlorobenzene | | 146 | 10.295 | 10.295 (1.039) | | 651646 | 25.0000 | 23.972 |
| 84 1,2-Dibromo-3-chloropropane | | 157 | 11.052 | 11.052 (1.116) | | 63184 | 25.0000 | 24.703 |
| 85 1,2,4-Trichlorobenzene | | 180 | 11.892 | 11.892 (1.201) | | 261752 | 25.0000 | 22.617 |
| 86 Hexachlorobutadiene | | 225 | 12.070 | 12.070 (1.219) | | 126348 | 25.0000 | 23.115 |
| 87 Naphthalene | | 128 | 12.129 | 12.129 (1.225) | | 610547 | 25.0000 | 22.352 |
| 88 1,2,3-Trichlorobenzene | | 180 | 12.377 | 12.377 (1.250) | | 182675 | 25.0000 | 22.011 |
| 98 Cyclohexane | | 56 | 4.568 | 4.568 (0.908) | | 334819 | 25.0000 | 19.770 |
| 143 Methyl Acetate | | 43 | 2.994 | 2.994 (0.595) | | 462866 | 50.0000 | 48.879 |
| 144 Methylcyclohexane | | 83 | 5.514 | 5.514 (1.096) | | 275331 | 25.0000 | 19.373 |
| 141 1,3,5-Trichlorobenzene | | 180 | 11.277 | 11.277 (1.139) | | 338982 | 25.0000 | 22.189 |

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Instrument: azux11.i
 Operator: 43562
 Column diameter: 0.18



STL North Canton

VOLATILE REPORT SW-846 Method
Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23874.D
Lab Smp Id: 10NG-IC
Inj Date : 14-SEP-2004 15:19
Operator : 43582 Inst ID: A3UX11.i
Smp Info : 10NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 2
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:47 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 6 Calibration Sample, Level: 2
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|----------------------------|-----------|---------|---------------|---------|---------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| * 1 Fluorobenzene | 96 | 5.029 | 5.029 (1.000) | 2293402 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1817387 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 990181 | 50.0000 | | |
| \$ 4 Dibromofluoromethane | 113 | 4.473 | 4.473 (0.889) | 106274 | 10.0000 | 9.953 | |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.745 | 4.745 (0.944) | 141757 | 10.0000 | 9.655 | |
| \$ 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 427652 | 10.0000 | 9.795 | |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 185658 | 10.0000 | 10.008 | |
| 8 Dichlorodifluoromethane | 85 | 1.526 | 1.526 (0.304) | 108953 | 10.0000 | 9.087 | |
| 9 Chloromethane | 50 | 1.680 | 1.680 (0.334) | 228871 | 10.0000 | 10.691 | |
| 10 Vinyl Chloride | 62 | 1.763 | 1.763 (0.351) | 146078 | 10.0000 | 9.976 | |
| 11 Bromomethane | 94 | 2.047 | 2.047 (0.407) | 75436 | 10.0000 | 10.861 | |
| 12 Chloroethane | 64 | 2.118 | 2.118 (0.421) | 113260 | 10.0000 | 10.524 | |
| 13 Trichlorofluoromethane | 101 | 2.307 | 2.307 (0.459) | 142867 | 10.0000 | 9.308 | |
| 15 Acrolein | 56 | 2.615 | 2.615 (0.520) | 140593 | 100.000 | 98.140 | |
| 16 Acetone | 43 | 2.722 | 2.722 (0.541) | 128042 | 20.0000 | 20.736 | |
| 17 1,1-Dichloroethene | 96 | 2.710 | 2.710 (0.539) | 105536 | 10.0000 | 10.290 | |
| 18 Freon-113 | 151 | 2.745 | 2.745 (0.546) | 76261 | 10.0000 | 11.396 | |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|---------------------------------|-----------|---------|---------------|--------|--------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| 19 Iodomethane | 142 | 2.840 | 2.840 (0.565) | | 159856 | 10.0000 | 10.545 |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 (0.577) | | 400515 | 10.0000 | 10.454 |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 (0.612) | | 251128 | 10.0000 | 10.978 |
| 22 Acetonitrile | 41 | 2.946 | 2.946 (0.586) | | 138464 | 100.000 | 101.73 |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 (0.647) | | 439101 | 100.000 | 103.74 |
| 24 Methyl tert-butyl ether | 73 | 3.301 | 3.301 (0.657) | | 308600 | 10.0000 | 9.385 |
| 25 trans-1,2-Dichloroethene | 96 | 3.313 | 3.313 (0.659) | | 126088 | 10.0000 | 10.363 |
| 26 Hexane | 86 | 3.538 | 3.538 (0.704) | | 22298 | 10.0000 | 11.260 |
| 27 Vinyl acetate | 43 | 3.656 | 3.656 (0.727) | | 175571 | 10.0000 | 9.018 |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 (0.722) | | 226350 | 10.0000 | 10.198 |
| 29 tert-Butyl Alcohol | 59 | 3.148 | 3.148 (0.626) | | 185158 | 200.000 | 207.57 |
| 30 2-Butanone | 43 | 4.082 | 4.082 (0.812) | | 117242 | 20.0000 | 18.664 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 261363 | 20.0000 | 20.945 |
| 32 cis-1,2-dichloroethene | 96 | 4.094 | 4.094 (0.814) | | 135275 | 10.0000 | 10.582 |
| 33 2,2-Dichloropropane | 77 | 4.106 | 4.106 (0.816) | | 141153 | 10.0000 | 10.553 |
| 34 Bromochloromethane | 128 | 4.283 | 4.283 (0.852) | | 65133 | 10.0000 | 11.003 |
| 35 Chloroform | 83 | 4.343 | 4.343 (0.864) | | 231979 | 10.0000 | 10.397 |
| 36 Tetrahydrofuran | 42 | 4.331 | 4.331 (0.861) | | 38288 | 10.0000 | 11.006 |
| 37 1,1,1-Trichloroethane | 97 | 4.508 | 4.508 (0.896) | | 171781 | 10.0000 | 10.194 |
| 38 1,1-Dichloropropene | 75 | 4.650 | 4.650 (0.925) | | 163880 | 10.0000 | 10.435 |
| 39 Carbon Tetrachloride | 117 | 4.650 | 4.650 (0.925) | | 134205 | 10.0000 | 10.143 |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 (0.958) | | 194488 | 10.0000 | 10.668 |
| 41 Benzene | 78 | 4.816 | 4.816 (0.958) | | 545878 | 10.0000 | 10.338 |
| 42 Trichloroethene | 130 | 5.337 | 5.337 (1.061) | | 121044 | 10.0000 | 10.045 |
| 43 1,2-Dichloropropane | 63 | 5.526 | 5.526 (1.099) | | 138162 | 10.0000 | 10.607 |
| 44 1,4-Dioxane | 88 | 5.621 | 5.621 (1.118) | | 53454 | 500.000 | 478.78 (A) |
| 45 Dibromomethane | 93 | 5.621 | 5.621 (1.118) | | 76444 | 10.0000 | 10.591 |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 (1.144) | | 176533 | 10.0000 | 10.303 |
| 47 2-Chloroethyl vinyl ether | 63 | 5.987 | 5.987 (1.191) | | 135749 | 20.0000 | 18.535 |
| 48 cis-1,3-Dichloropropene | 75 | 6.129 | 6.129 (1.219) | | 218946 | 10.0000 | 10.444 |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 (1.242) | | 216822 | 20.0000 | 19.234 |
| 50 Toluene | 91 | 6.437 | 6.437 (0.838) | | 563016 | 10.0000 | 10.373 |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 (0.860) | | 193073 | 10.0000 | 9.921 |
| 52 Ethyl Methacrylate | 69 | 6.674 | 6.674 (0.869) | | 163654 | 10.0000 | 9.774 |
| 53 1,1,2-Trichloroethane | 97 | 6.768 | 6.768 (0.881) | | 114065 | 10.0000 | 10.360 |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 (0.901) | | 215113 | 10.0000 | 10.370 |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 (0.903) | | 95253 | 10.0000 | 10.832 |
| 56 2-Hexanone | 43 | 6.981 | 6.981 (0.909) | | 173068 | 20.0000 | 19.692 |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 (0.929) | | 121301 | 10.0000 | 10.330 |
| 58 1,2-Dibromoethane | 107 | 7.254 | 7.254 (0.945) | | 114994 | 10.0000 | 10.601 |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 (1.003) | | 377942 | 10.0000 | 10.754 |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 (1.012) | | 125087 | 10.0000 | 10.245 |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 (1.015) | | 184529 | 10.0000 | 10.397 |
| 62 m + p-Xylene | 106 | 7.904 | 7.904 (1.029) | | 470158 | 20.0000 | 20.568 |
| M 63 Xylenes (total) | 106 | | | | 701783 | 30.0000 | 30.890 |
| 64 Xylene-o | 106 | 8.283 | 8.283 (1.079) | | 231625 | 10.0000 | 10.322 |
| 65 Styrene | 104 | 8.295 | 8.295 (1.080) | | 406292 | 10.0000 | 10.124 |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|--------------------------------|-----------|---------|--------|----------------|--------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| 66 Bromoform | ==== | 173 | 8.472 | 8.472 (1.103) | | 81750 | 10.0000 | 10.242 |
| 67 Isopropylbenzene | ==== | 105 | 8.626 | 8.626 (1.123) | | 495185 | 10.0000 | 9.908 |
| 68 1,1,2,2-Tetrachloroethane | ==== | 83 | 8.898 | 8.898 (0.898) | | 154970 | 10.0000 | 10.382 |
| 69 1,4-Dichloro-2-butene | ==== | 53 | 8.957 | 8.957 (0.904) | | 47422 | 10.0000 | 9.656 |
| 70 1,2,3-Trichloropropane | ==== | 110 | 8.946 | 8.946 (0.903) | | 49915 | 10.0000 | 10.191 |
| 71 Bromobenzene | ==== | 156 | 8.934 | 8.934 (0.902) | | 154286 | 10.0000 | 10.444 |
| 72 n-Propylbenzene | ==== | 120 | 9.028 | 9.028 (0.912) | | 140117 | 10.0000 | 10.173 |
| 73 2-Chlorotoluene | ==== | 126 | 9.111 | 9.111 (0.920) | | 137109 | 10.0000 | 10.052 |
| 74 1,3,5-Trimethylbenzene | ==== | 105 | 9.194 | 9.194 (0.928) | | 449094 | 10.0000 | 9.798 |
| 75 4-Chlorotoluene | ==== | 126 | 9.218 | 9.218 (0.931) | | 150151 | 10.0000 | 10.304 |
| 76 tert-Butylbenzene | ==== | 119 | 9.514 | 9.514 (0.961) | | 384632 | 10.0000 | 10.359 |
| 77 1,2,4-Trimethylbenzene | ==== | 105 | 9.561 | 9.561 (0.965) | | 494544 | 10.0000 | 10.108 |
| 78 sec-Butylbenzene | ==== | 105 | 9.727 | 9.727 (0.982) | | 506228 | 10.0000 | 10.072 |
| 79 4-Isopropyltoluene | ==== | 119 | 9.869 | 9.869 (0.996) | | 417405 | 10.0000 | 9.982 |
| 80 1,3-Dichlorobenzene | ==== | 146 | 9.845 | 9.845 (0.994) | | 280475 | 10.0000 | 10.284 |
| 81 1,4-Dichlorobenzene | ==== | 146 | 9.928 | 9.928 (1.002) | | 298356 | 10.0000 | 10.456 |
| 82 n-Butylbenzene | ==== | 91 | 10.271 | 10.271 (1.037) | | 380534 | 10.0000 | 10.187 |
| 83 1,2-Dichlorobenzene | ==== | 146 | 10.295 | 10.295 (1.039) | | 278506 | 10.0000 | 10.368 |
| 84 1,2-Dibromo-3-chloropropane | ==== | 157 | 11.064 | 11.064 (1.117) | | 24938 | 10.0000 | 9.866 |
| 85 1,2,4-Trichlorobenzene | ==== | 180 | 11.892 | 11.892 (1.201) | | 120755 | 10.0000 | 10.559 |
| 86 Hexachlorobutadiene | ==== | 225 | 12.070 | 12.070 (1.219) | | 62649 | 10.0000 | 9.935 |
| 87 Naphthalene | ==== | 128 | 12.129 | 12.129 (1.225) | | 251285 | 10.0000 | 9.310 |
| 88 1,2,3-Trichlorobenzene | ==== | 180 | 12.377 | 12.377 (1.250) | | 87516 | 10.0000 | 10.671 |
| 98 Cyclohexane | ==== | 56 | 4.567 | 4.567 (0.908) | | 177719 | 10.0000 | 10.533 |
| 143 Methyl Acetate | ==== | 43 | 2.994 | 2.994 (0.595) | | 199578 | 20.0000 | 21.155 |
| 144 Methylcyclohexane | ==== | 83 | 5.514 | 5.514 (1.096) | | 144963 | 10.0000 | 10.239 |
| 141 1,3,5-Trichlorobenzene | ==== | 180 | 11.277 | 11.277 (1.139) | | 157787 | 10.0000 | 10.452 |

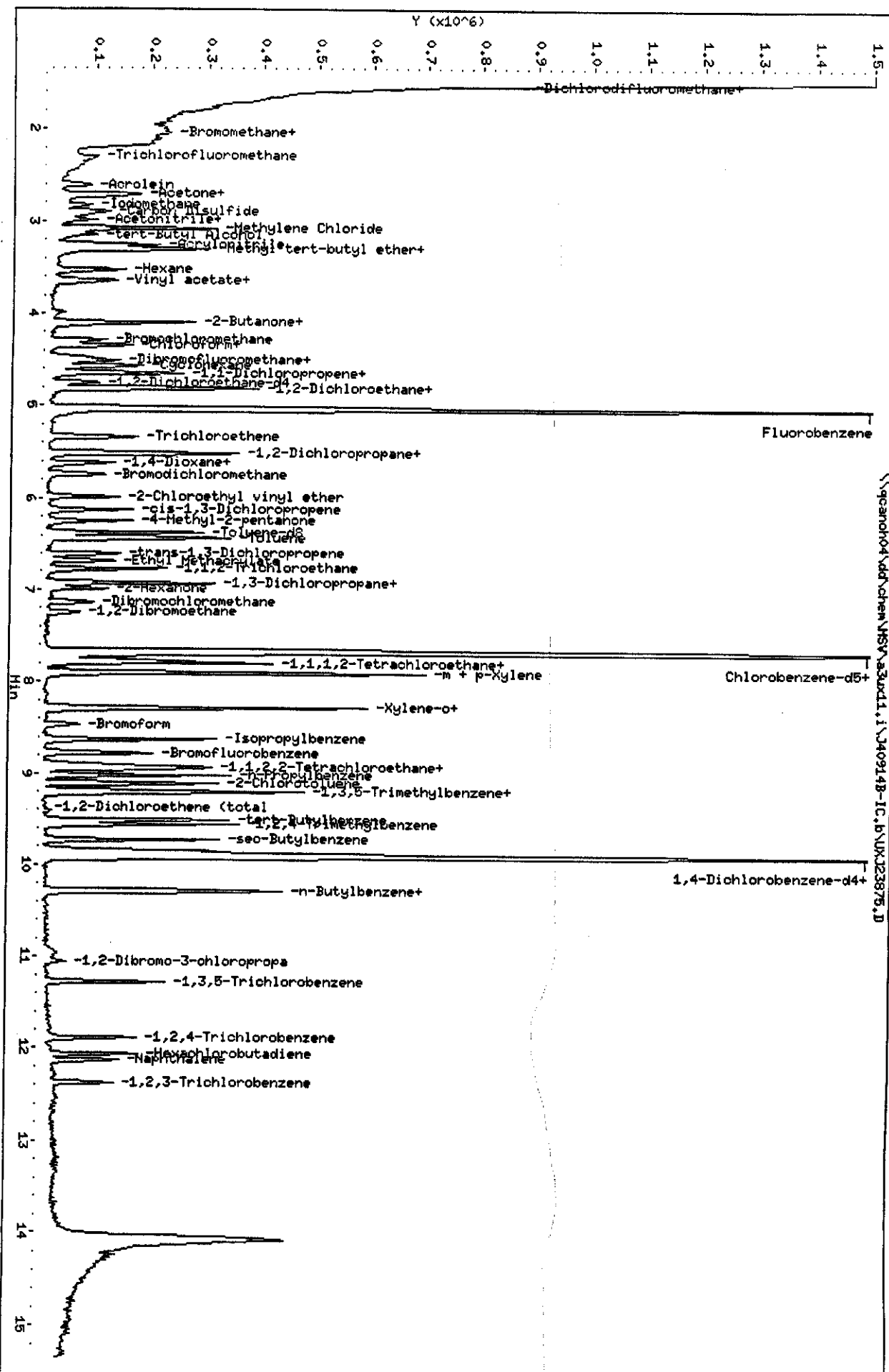
QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\qcanoh04\\dd\\chem\\HSV\\a3x11.i\\J40914B-IC.b\\UXJ23875.D
Date : 14-SEP-2004 15:41
Client ID:
Sample Info: SUG-IC
Purge Volume: 5.0
Column Phase: DB624

Instrument: a3x11.i

Operator: 43582
Column diameter: 0.18
\\qcanoh04\\dd\\chem\\HSV\\a3x11.i\\J40914B-IC.b\\UXJ23875.D



STL North Canton

VOLATILE REPORT SW-846 Method
Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23875.D
Lab Smp Id: SNG-IC
Inj Date : 14-SEP-2004 15:41
Operator : 43582 Inst ID: a3ux11.i
Smp Info : SNG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 1
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:47 evans1 Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 7 Calibration Sample, Level: 1
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|----------------------------|-----------|---------|-------|---------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * 1 Fluorobenzene | 96 | 5.029 | 5.029 | (1.000) | 2251983 | 50.0000 | | |
| * 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 | (1.000) | 1826599 | 50.0000 | | |
| * 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 | (1.000) | 937868 | 50.0000 | | |
| \$ 4 Dibromofluoromethane | 113 | 4.473 | 4.473 | (0.889) | 51569 | 5.00000 | 4.918 | |
| \$ 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 | (0.946) | 70414 | 5.00000 | 4.884 | |
| \$ 6 Toluene-d8 | 98 | 6.378 | 6.378 | (0.831) | 210171 | 5.00000 | 4.789 | |
| \$ 7 Bromofluorobenzene | 95 | 8.780 | 8.780 | (1.143) | 88368 | 5.00000 | 4.739 | |
| 8 Dichlorodifluoromethane | 85 | 1.526 | 1.526 | (0.304) | 64938 | 5.00000 | 5.516 | |
| 9 Chloromethane | 50 | 1.668 | 1.668 | (0.332) | 125378 | 5.00000 | 5.964 | |
| 10 Vinyl Chloride | 62 | 1.763 | 1.763 | (0.351) | 72850 | 5.00000 | 5.066 | |
| 11 Bromomethane | 94 | 2.035 | 2.035 | (0.405) | 38866 | 5.00000 | 5.699 | |
| 12 Chloroethane | 64 | 2.118 | 2.118 | (0.421) | 52345 | 5.00000 | 4.953 | |
| 13 Trichlorofluoromethane | 101 | 2.307 | 2.307 | (0.459) | 86493 | 5.00000 | 5.739 | |
| 15 Acrolein | 56 | 2.615 | 2.615 | (0.520) | 72041 | 50.0000 | 51.213 | |
| 16 Acetone | 43 | 2.722 | 2.722 | (0.541) | 72580 | 10.0000 | 9.714 | |
| 17 1,1-Dichloroethene | 96 | 2.710 | 2.710 | (0.539) | 55428 | 5.00000 | 5.504 | |
| 18 Freon-113 | 151 | 2.745 | 2.745 | (0.546) | 37991 | 5.00000 | 5.276 | |

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|---------------------------------|-----------|---------|-------|---------|--------|----------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| 19 Iodomethane | 142 | 2.828 | 2.828 | (0.562) | 70148 | 5.00000 | 4.712 |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 | (0.577) | 205480 | 5.00000 | 5.462 |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 | (0.612) | 169409 | 5.00000 | 4.472 |
| 22 Acetonitrile | 41 | 2.946 | 2.946 | (0.586) | 74099 | 50.0000 | 55.440 |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 | (0.647) | 205768 | 50.0000 | 49.508 |
| 24 Methyl tert-butyl ether | 73 | 3.301 | 3.301 | (0.657) | 147953 | 5.00000 | 4.582 |
| 25 trans-1,2-Dichloroethene | 96 | 3.313 | 3.313 | (0.659) | 63645 | 5.00000 | 5.327 |
| 26 Hexane | 86 | 3.526 | 3.526 | (0.701) | 11663 | 5.00000 | 5.789 |
| 27 Vinyl acetate | 43 | 3.656 | 3.656 | (0.727) | 93501 | 5.00000 | 4.891 |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 | (0.722) | 115670 | 5.00000 | 5.307 |
| 29 tert-Butyl Alcohol | 59 | 3.148 | 3.148 | (0.626) | 92424 | 100.000 | 105.52 |
| 30 2-Butanone | 43 | 4.094 | 4.094 | (0.814) | 71588 | 10.0000 | 11.606 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 128452 | 10.0000 | 10.490 |
| 32 cis-1,2-dichloroethene | 96 | 4.106 | 4.106 | (0.816) | 64807 | 5.00000 | 5.163 |
| 33 2,2-Dichloropropane | 77 | 4.106 | 4.106 | (0.816) | 66643 | 5.00000 | 5.074 |
| 34 Bromochloromethane | 128 | 4.295 | 4.295 | (0.854) | 28755 | 5.00000 | 4.947 |
| 35 Chloroform | 83 | 4.343 | 4.343 | (0.864) | 114878 | 5.00000 | 5.243 |
| 36 Tetrahydrofuran | 42 | 4.331 | 4.331 | (0.861) | 22560 | 5.00000 | 6.678 |
| 37 1,1,1-Trichloroethane | 97 | 4.508 | 4.508 | (0.896) | 88473 | 5.00000 | 5.347 |
| 38 1,1-Dichloropropene | 75 | 4.650 | 4.650 | (0.925) | 78641 | 5.00000 | 5.099 |
| 39 Carbon Tetrachloride | 117 | 4.650 | 4.650 | (0.925) | 68204 | 5.00000 | 5.250 |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 | (0.958) | 90135 | 5.00000 | 5.035 |
| 41 Benzene | 78 | 4.816 | 4.816 | (0.958) | 283739 | 5.00000 | 5.472 |
| 42 Trichloroethene | 130 | 5.349 | 5.349 | (1.064) | 61441 | 5.00000 | 5.192 |
| 43 1,2-Dichloropropane | 63 | 5.514 | 5.514 | (1.096) | 66815 | 5.00000 | 5.224 |
| 44 1,4-Dioxane | 88 | 5.621 | 5.621 | (1.118) | 27287 | 250.000 | 248.90 (A) |
| 45 Dibromomethane | 93 | 5.621 | 5.621 | (1.118) | 34422 | 5.00000 | 4.857 |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 | (1.144) | 86009 | 5.00000 | 5.112 |
| 47 2-Chloroethyl vinyl ether | 63 | 5.988 | 5.988 | (1.191) | 62831 | 10.0000 | 8.737 |
| 48 cis-1,3-Dichloropropene | 75 | 6.129 | 6.129 | (1.219) | 97774 | 5.00000 | 4.750 |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 | (1.242) | 106568 | 10.0000 | 9.627 |
| 50 Toluene | 91 | 6.437 | 6.437 | (0.838) | 271647 | 5.00000 | 4.980 |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 | (0.860) | 92511 | 5.00000 | 4.730 |
| 52 Ethyl Methacrylate | 69 | 6.686 | 6.686 | (0.871) | 71117 | 5.00000 | 4.226 |
| 53 1,1,2-Trichloroethane | 97 | 6.780 | 6.780 | (0.883) | 55141 | 5.00000 | 4.983 |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 | (0.901) | 104493 | 5.00000 | 5.012 |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 | (0.903) | 46594 | 5.00000 | 5.272 |
| 56 2-Hexanone | 43 | 6.993 | 6.993 | (0.911) | 81226 | 10.0000 | 9.196 |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 | (0.929) | 56564 | 5.00000 | 4.793 |
| 58 1,2-Dibromoethane | 107 | 7.242 | 7.242 | (0.943) | 49174 | 5.00000 | 4.510 |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 | (1.003) | 176312 | 5.00000 | 4.992 |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 | (1.012) | 61700 | 5.00000 | 5.028 |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 | (1.015) | 84068 | 5.00000 | 4.713 |
| 62 m + p-Xylene | 106 | 7.904 | 7.904 | (1.029) | 221783 | 10.0000 | 9.654 |
| M 63 Xylenes (total) | 106 | | | | 325206 | 15.0000 | 14.239 |
| 64 Xylene-o | 106 | 8.283 | 8.283 | (1.079) | 103423 | 5.00000 | 4.586 |
| 65 Styrene | 104 | 8.295 | 8.295 | (1.080) | 183225 | 5.00000 | 4.543 |

| Compounds | QUANT SIG | MASS | RT | EXP RT | REL RT | RESPONSE | AMOUNTS | |
|--------------------------------|-----------|------|--------|----------------|--------|----------|------------------|-----------------|
| | | | | | | | CAL-AMT (ng) | ON-COL (ng) |
| 66 Bromoform | ==== | 173 | 8.472 | 8.472 (1.103) | | 36774 | 5.00000 | 4.584 |
| 67 Isopropylbenzene | ==== | 105 | 8.626 | 8.626 (1.123) | | 245674 | 5.00000 | 4.891 |
| 68 1,1,2,2-Tetrachloroethane | ==== | 83 | 8.898 | 8.898 (0.898) | | 69472 | 5.00000 | 4.914 |
| 69 1,4-Dichloro-2-butene | ==== | 53 | 8.958 | 8.958 (0.904) | | 21722 | 5.00000 | 4.670 |
| 70 1,2,3-Trichloropropane | ==== | 110 | 8.946 | 8.946 (0.903) | | 23727 | 5.00000 | 5.114 |
| 71 Bromobenzene | ==== | 156 | 8.934 | 8.934 (0.902) | | 69868 | 5.00000 | 4.993 |
| 72 n-Propylbenzene | ==== | 120 | 9.029 | 9.029 (0.912) | | 67269 | 5.00000 | 5.156 |
| 73 2-Chlorotoluene | ==== | 126 | 9.111 | 9.111 (0.920) | | 67172 | 5.00000 | 5.199 |
| 74 1,3,5-Trimethylbenzene | ==== | 105 | 9.194 | 9.194 (0.928) | | 211928 | 5.00000 | 4.882 |
| 75 4-Chlorotoluene | ==== | 126 | 9.218 | 9.218 (0.931) | | 71891 | 5.00000 | 5.209 |
| 76 tert-Butylbenzene | ==== | 119 | 9.514 | 9.514 (0.961) | | 177303 | 5.00000 | 5.041 |
| 77 1,2,4-Trimethylbenzene | ==== | 105 | 9.561 | 9.561 (0.965) | | 218676 | 5.00000 | 4.719 |
| 78 sec-Butylbenzene | ==== | 105 | 9.727 | 9.727 (0.982) | | 249496 | 5.00000 | 5.241 |
| 79 4-Isopropyltoluene | ==== | 119 | 9.869 | 9.869 (0.996) | | 190215 | 5.00000 | 4.803 |
| 80 1,3-Dichlorobenzene | ==== | 146 | 9.845 | 9.845 (0.994) | | 141050 | 5.00000 | 5.460 |
| 81 1,4-Dichlorobenzene | ==== | 146 | 9.928 | 9.928 (1.002) | | 139945 | 5.00000 | 5.178 |
| 82 n-Butylbenzene | ==== | 91 | 10.271 | 10.271 (1.037) | | 174632 | 5.00000 | 4.936 |
| 83 1,2-Dichlorobenzene | ==== | 146 | 10.295 | 10.295 (1.039) | | 135942 | 5.00000 | 5.343 |
| 84 1,2-Dibromo-3-chloropropane | ==== | 157 | 11.064 | 11.064 (1.117) | | 11856 | 5.00000 | 4.952 |
| 85 1,2,4-Trichlorobenzene | ==== | 180 | 11.892 | 11.892 (1.201) | | 52887 | 5.00000 | 4.882 |
| 86 Hexachlorobutadiene | ==== | 225 | 12.070 | 12.070 (1.219) | | 36648 | 5.00000 | 4.872 |
| 87 Naphthalene | ==== | 128 | 12.129 | 12.129 (1.225) | | 112143 | 5.00000 | 4.386 |
| 88 1,2,3-Trichlorobenzene | ==== | 180 | 12.377 | 12.377 (1.250) | | 35937 | 5.00000 | 4.626 |
| 98 Cyclohexane | ==== | 56 | 4.568 | 4.568 (0.908) | | 84417 | 5.00000 | 5.095 |
| 143 Methyl Acetate | ==== | 43 | 2.994 | 2.994 (0.595) | | 100499 | 10.0000 | 10.849 |
| 144 Methylcyclohexane | ==== | 83 | 5.514 | 5.514 (1.096) | | 78015 | 5.00000 | 5.612 |
| 141 1,3,5-Trichlorobenzene | ==== | 180 | 11.277 | 11.277 (1.139) | | 80418 | 5.00000 | 5.624 |

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

STL North Canton

RECOVERY REPORT

Client Name:
 Sample Matrix: LIQUID
 Lab Smp Id: ICV
 Level: LOW
 Data Type: MS DATA
 SpikeList File: plexus-ck.spk
 Sublist File: 2-8260.SUB
 Method File: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
 Misc Info: J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 3

Client SDG: SDGa00594
 Fraction: VOA
 Operator: 43582
 SampleType: METHSPIKE
 Quant Type: ISTD

| SPIKE COMPOUND | CONC ADDED ug/L | CONC RECOVERED ug/L | % RECOVERED | LIMITS |
|-------------------------|-----------------|---------------------|-------------|--------|
| 17 1,1-Dichloroethene | 10.000 | 9.507 | 95.07 | 45-155 |
| 42 Trichloroethene | 10.000 | 9.342 | 93.42 | 45-155 |
| 59 Chlorobenzene | 10.000 | 9.443 | 94.43 | 45-155 |
| 50 Toluene | 10.000 | 9.460 | 94.60 | 45-155 |
| 41 Benzene | 10.000 | 9.294 | 92.94 | 45-155 |
| 16 Acetone | 10.000 | 6.134 | 61.34 | 45-155 |
| 20 Carbon Disulfide | 10.000 | 9.433 | 94.33 | 45-155 |
| 9 Chloromethane | 10.000 | 8.196 | 81.96 | 45-155 |
| 11 Bromomethane | 10.000 | 8.864 | 88.64 | 45-155 |
| 10 Vinyl Chloride | 10.000 | 8.534 | 85.34 | 45-155 |
| 12 Chloroethane | 10.000 | 8.736 | 87.36 | 45-155 |
| 21 Methylene Chloride | 10.000 | 9.642 | 96.42 | 45-155 |
| 28 1,1-Dichloroethane | 10.000 | 9.526 | 95.26 | 45-155 |
| M 31 1,2-Dichloroethene | 20.000 | 18.957 | 94.78 | 45-155 |
| 35 Chloroform | 10.000 | 9.419 | 94.19 | 45-155 |
| 40 1,2-Dichloroethane | 10.000 | 9.839 | 98.39 | 45-155 |
| 30 2-Butanone | 10.000 | 8.378 | 83.78 | 45-155 |
| 37 1,1,1-Trichloroeth | 10.000 | 9.100 | 91.00 | 45-155 |
| 39 Carbon Tetrachlori | 10.000 | 9.159 | 91.59 | 45-155 |
| 46 Bromodichlorometha | 10.000 | 9.513 | 95.13 | 45-155 |
| 43 1,2-Dichloropropan | 10.000 | 9.463 | 94.63 | 45-155 |
| 48 cis-1,3-Dichloropr | 10.000 | 9.690 | 96.90 | 45-155 |
| 57 Dibromochlorometha | 10.000 | 10.024 | 100.24 | 45-155 |
| 53 1,1,2-Trichloroeth | 10.000 | 9.656 | 96.56 | 45-155 |
| 51 trans-1,3-Dichloro | 10.000 | 9.392 | 93.92 | 45-155 |
| 66 Bromoform | 10.000 | 9.919 | 99.19 | 45-155 |
| 49 4-Methyl-2-pentano | 10.000 | 9.709 | 97.09 | 45-155 |
| 56 2-Hexanone | 10.000 | 8.736 | 87.36 | 45-155 |
| 55 Tetrachloroethene | 10.000 | 9.092 | 90.92 | 45-155 |
| 68 1,1,2,2-Tetrachlor | 10.000 | 10.303 | 103.03 | 45-155 |
| 61 Ethylbenzene | 10.000 | 9.427 | 94.27 | 45-155 |
| 65 Styrene | 10.000 | 9.617 | 96.17 | 45-155 |
| M 63 Xylenes (total) | 30.000 | 28.176 | 93.92 | 45-155 |

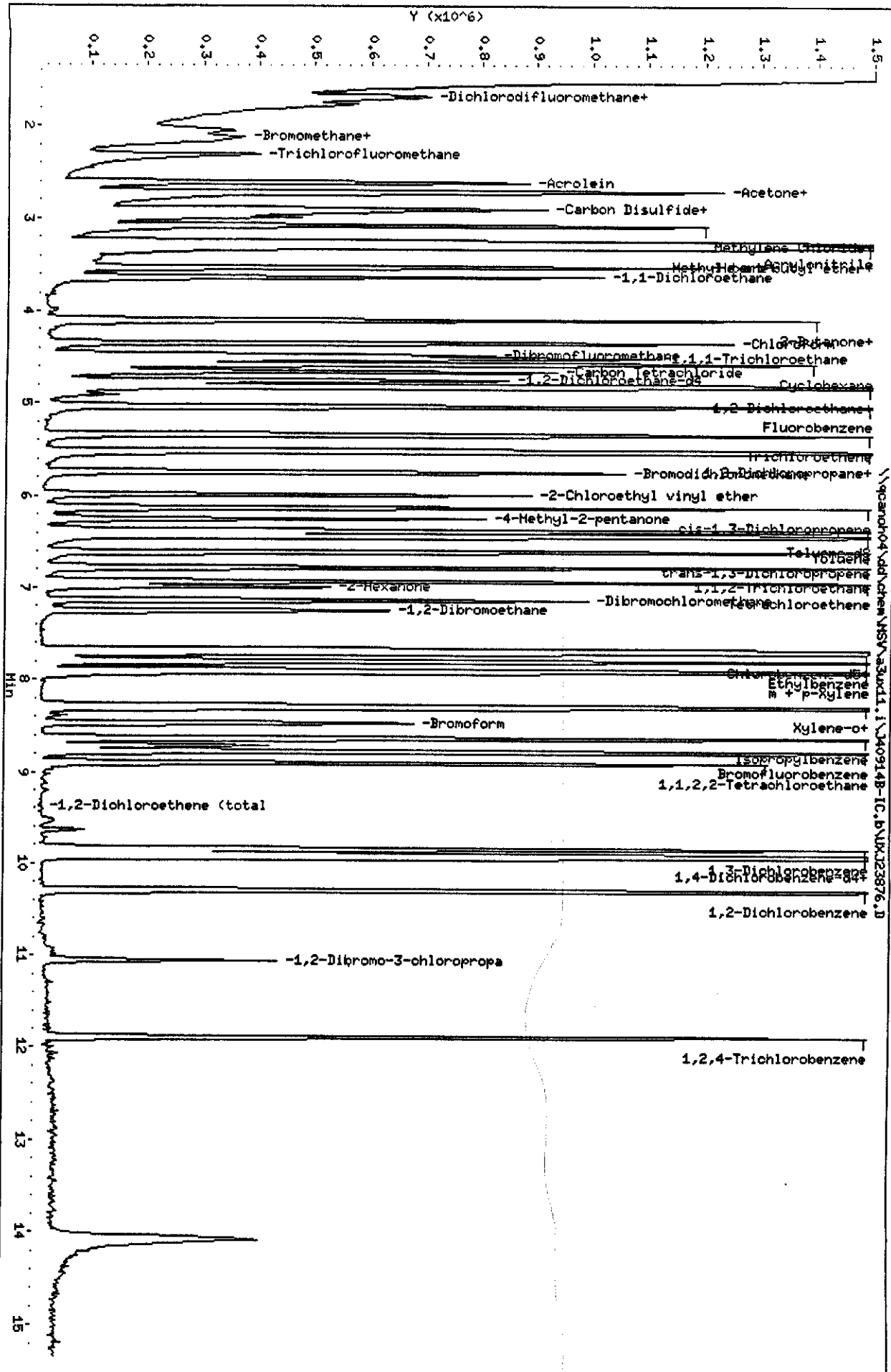
| SPIKE COMPOUND | CONC ADDED ug/L | CONC RECOVERED ug/L | % RECOVERED | LIMITS |
|-----------------------|-----------------|---------------------|-------------|--------|
| 32 cis-1,2-dichloroet | 10.000 | 9.524 | 95.24 | 45-155 |
| 25 trans-1,2-Dichloro | 10.000 | 9.432 | 94.32 | 45-155 |
| 8 Dichlorodifluorome | 10.000 | 9.117 | 91.17 | 45-155 |
| 13 Trichlorofluoromet | 10.000 | 9.634 | 96.34 | 45-155 |
| 18 Freon-113 | 10.000 | 11.317 | 113.17 | 45-155 |
| 24 Methyl tert-butyl | 10.000 | 9.278 | 92.78 | 45-155 |
| 58 1,2-Dibromoethane | 10.000 | 9.902 | 99.02 | 45-155 |
| 67 Isopropylbenzene | 10.000 | 9.750 | 97.50 | 45-155 |
| 80 1,3-Dichlorobenzen | 10.000 | 9.577 | 95.77 | 45-155 |
| 81 1,4-Dichlorobenzen | 10.000 | 9.879 | 98.79 | 45-155 |
| 83 1,2-Dichlorobenzen | 10.000 | 9.570 | 95.70 | 45-155 |
| 84 1,2-Dibromo-3-chlo | 10.000 | 10.335 | 103.35 | 45-155 |
| 85 1,2,4-Trichloroben | 10.000 | 9.534 | 95.34 | 45-155 |
| 98 Cyclohexane | 10.000 | 9.159 | 91.59 | 45-155 |
| 143 Methyl Acetate | 10.000 | 9.302 | 93.02 | 45-155 |
| 144 Methylcyclohexane | 10.000 | 8.960 | 89.60 | 45-155 |

| SURROGATE COMPOUND | CONC ADDED ug/L | CONC RECOVERED ug/L | % RECOVERED | LIMITS |
|-------------------------|-----------------|---------------------|-------------|--------|
| \$ 4 Dibromofluorometha | 10.000 | 9.507 | 95.07 | 73-122 |
| \$ 5 1,2-Dichloroethane | 10.000 | 9.840 | 98.40 | 61-128 |
| \$ 6 Toluene-d8 | 10.000 | 9.847 | 98.47 | 76-110 |
| \$ 7 Bromofluorobenzene | 10.000 | 10.152 | 101.52 | 74-116 |

Data File: \\qcanch04\\dd\\chem\\MSV\\a30x11.i\\J40914B-IC.b\\UKJ23876.D
Date : 14-SEP-2004 16:04
Client ID:
Sample Info: ICQ
Purge Volume: 5.0
Column Phase: DB624

Instrument: a30x11.i

Operator: 43582
Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23876.D
Report Date: 15-Sep-2004 13:02

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23876.D
Lab Smp Id: ICV
Inj Date : 14-SEP-2004 16:04
Operator : 43582 Inst ID: a3ux11.i
Smp Info : ICV
Misc Info : J40914B-IC,8260LLUX11,2-8260.SUB,43582,3
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 13:00 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 8 QC Sample: METHSPIKE
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|-----------|--------------------------|----------------|------------------------|---------------|---------|----------|---------------|--|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) (ug/L) | |
| * | 1 Fluorobenzene | 96 | 5.029 | 5.029 (1.000) | 2348288 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1882362 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 979729 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.473 | 4.473 (0.889) | 519715 | 47.5349 | 9.507 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.745 | 4.757 (0.944) | 739678 | 49.1997 | 9.840 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 2226509 | 49.2353 | 9.847 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 975336 | 50.7609 | 10.152 | |
| 8 | Dichlorodifluoromethane | 85 | 1.527 | 1.526 (0.304) | 559620 | 45.5836 | 9.117 | |
| 9 | Chloromethane | 50 | 1.680 | 1.668 (0.334) | 898254 | 40.9791 | 8.196 | |
| 10 | Vinyl Chloride | 62 | 1.775 | 1.763 (0.353) | 639778 | 42.6691 | 8.534 | |
| 11 | Bromomethane | 94 | 2.047 | 2.035 (0.407) | 315198 | 44.3200 | 8.864 | |
| 12 | Chloroethane | 64 | 2.118 | 2.118 (0.421) | 481308 | 43.6780 | 8.736 | |
| 13 | Trichlorofluoromethane | 101 | 2.308 | 2.307 (0.459) | 756987 | 48.1676 | 9.634 | |
| 15 | Acrolein | 56 | 2.615 | 2.615 (0.520) | 1028164 | 700.930 | 140.18 | |
| 16 | Acetone | 43 | 2.722 | 2.722 (0.541) | 180849 | 30.6678 | 6.134 | |
| 17 | 1,1-Dichloroethene | 96 | 2.710 | 2.710 (0.539) | 499203 | 47.5355 | 9.507 | |
| 18 | Freon-113 | 151 | 2.734 | 2.745 (0.544) | 373218 | 56.5832 | 11.317 | |
| 19 | Iodomethane | 142 | Compound Not Detected. | | | | | |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | |
|---------------------------------|-----------|----------------|------------------------|---------|---------|----------|----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) |
| 20 Carbon Disulfide | 76 | 2.899 | 2.899 (0.577) | 1850176 | 47.1646 | 9.433 | |
| 21 Methylene Chloride | 84 | 3.077 | 3.077 (0.612) | 719449 | 48.2118 | 9.642 | |
| 22 Acetonitrile | 41 | 2.935 | 2.946 (0.584) | 665528 | 477.523 | 95.505 | |
| 23 Acrylonitrile | 53 | 3.254 | 3.254 (0.647) | 2230444 | 514.640 | 102.93 | |
| 24 Methyl tert-butyl ether | 73 | 3.302 | 3.301 (0.657) | 1561889 | 46.3911 | 9.278 | |
| 25 trans-1,2-Dichloroethene | 96 | 3.313 | 3.313 (0.659) | 587539 | 47.1615 | 9.432 | |
| 26 Hexane | 86 | 3.526 | 3.526 (0.701) | 107938 | 53.4083 | 10.682 | |
| 27 Vinyl acetate | 43 | 3.526 | 3.656 (0.701) | 387039 | 19.4150 | 3.883 | |
| 28 1,1-Dichloroethane | 63 | 3.633 | 3.633 (0.722) | 1082468 | 47.6309 | 9.526 | |
| 29 tert-Butyl Alcohol | 59 | 2.994 | 3.148 (0.595) | 39015 | 42.7151 | 8.543 | |
| 30 2-Butanone | 43 | 4.083 | 4.094 (0.812) | 269449 | 41.8924 | 8.378 | |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 1210887 | 94.7835 | 18.957 |
| 32 cis-1,2-dichloroethene | 96 | 4.094 | 4.106 (0.814) | 623348 | 47.6220 | 9.524 | |
| 33 2,2-Dichloropropane | 77 | | Compound Not Detected. | | | | |
| 34 Bromochloromethane | 128 | | Compound Not Detected. | | | | |
| 35 Chloroform | 83 | 4.343 | 4.343 (0.864) | 1075879 | 47.0934 | 9.419 | |
| 36 Tetrahydrofuran | 42 | 4.083 | 4.331 (0.812) | 19114 | 5.46454 | 1.093 | |
| 37 1,1,1-Trichloroethane | 97 | 4.509 | 4.508 (0.896) | 785098 | 45.5027 | 9.100 | |
| 38 1,1-Dichloropropene | 75 | | Compound Not Detected. | | | | |
| 39 Carbon Tetrachloride | 117 | 4.662 | 4.650 (0.927) | 620420 | 45.7962 | 9.159 | |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 (0.958) | 918291 | 49.1951 | 9.839 | |
| 41 Benzene | 78 | 4.816 | 4.816 (0.958) | 2512537 | 46.4726 | 9.294 | |
| 42 Trichloroethene | 130 | 5.337 | 5.349 (1.061) | 576326 | 46.7078 | 9.342 | |
| 43 1,2-Dichloropropane | 63 | 5.526 | 5.514 (1.099) | 631042 | 47.3158 | 9.463 | |
| 44 1,4-Dioxane | 88 | | Compound Not Detected. | | | | |
| 45 Dibromomethane | 93 | | Compound Not Detected. | | | | |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 (1.144) | 834464 | 47.5633 | 9.513 | |
| 47 2-Chloroethyl vinyl ether | 63 | 5.988 | 5.988 (1.191) | 393567 | 52.4826 | 10.496 | |
| 48 cis-1,3-Dichloropropene | 75 | 6.130 | 6.129 (1.219) | 1040015 | 48.4491 | 9.690 | |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 (1.242) | 560341 | 48.5456 | 9.709 | |
| 50 Toluene | 91 | 6.437 | 6.437 (0.838) | 2659243 | 47.3027 | 9.460 | |
| 51 trans-1,3-Dichloropropene | 75 | 6.603 | 6.603 (0.860) | 946487 | 46.9582 | 9.392 | |
| 52 Ethyl Methacrylate | 69 | | Compound Not Detected. | | | | |
| 53 1,1,2-Trichloroethane | 97 | 6.769 | 6.780 (0.881) | 550573 | 48.2825 | 9.656 | |
| 54 1,3-Dichloropropane | 76 | | Compound Not Detected. | | | | |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 (0.903) | 414030 | 45.4582 | 9.092 | |
| 56 2-Hexanone | 43 | 6.982 | 6.993 (0.909) | 397634 | 43.6825 | 8.736 | |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 (0.929) | 609608 | 50.1213 | 10.024 | |
| 58 1,2-Dibromoethane | 107 | 7.254 | 7.242 (0.945) | 556274 | 49.5113 | 9.902 | |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 (1.003) | 1718695 | 47.2166 | 9.443 | |
| 60 1,1,1,2-Tetrachloroethane | 131 | | Compound Not Detected. | | | | |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 (1.015) | 866451 | 47.1356 | 9.427 | |
| 62 m + p-Xylene | 106 | 7.905 | 7.904 (1.029) | 2209377 | 93.3186 | 18.664 | |
| M 63 Xylenes (total) | 106 | | | 3314793 | 140.878 | 28.176 | |
| 64 Xylene-o | 106 | 8.283 | 8.283 (1.079) | 1105416 | 47.5597 | 9.512 | |
| 65 Styrene | 104 | 8.295 | 8.295 (1.080) | 1998604 | 48.0849 | 9.617 | |
| 66 Bromoform | 173 | 8.473 | 8.472 (1.103) | 410026 | 49.5965 | 9.919 | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23876.D
 Report Date: 15-Sep-2004 13:02

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | |
|--------------------------------|-----------|------------------------|--------|---------|---------|----------|-------------------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 67 Isopropylbenzene | 105 | 8.626 | 8.626 | (1.123) | 2523492 | 48.7480 | 9.750 |
| 68 1,1,2,2-Tetrachloroethane | 83 | 8.899 | 8.898 | (0.898) | 760843 | 51.5174 | 10.303 |
| 69 1,4-Dichloro-2-butene | 53 | Compound Not Detected. | | | | | |
| 70 1,2,3-Trichloropropane | 110 | Compound Not Detected. | | | | | |
| 71 Bromobenzene | 156 | Compound Not Detected. | | | | | |
| 72 n-Propylbenzene | 120 | Compound Not Detected. | | | | | |
| 73 2-Chlorotoluene | 126 | Compound Not Detected. | | | | | |
| 74 1,3,5-Trimethylbenzene | 105 | Compound Not Detected. | | | | | |
| 75 4-Chlorotoluene | 126 | Compound Not Detected. | | | | | |
| 76 tert-Butylbenzene | 119 | Compound Not Detected. | | | | | |
| 77 1,2,4-Trimethylbenzene | 105 | Compound Not Detected. | | | | | |
| 78 sec-Butylbenzene | 105 | Compound Not Detected. | | | | | |
| 79 4-Isopropyltoluene | 119 | Compound Not Detected. | | | | | |
| 80 1,3-Dichlorobenzene | 146 | 9.845 | 9.845 | (0.994) | 1292290 | 47.8872 | 9.577 |
| 81 1,4-Dichlorobenzene | 146 | 9.928 | 9.928 | (1.002) | 1394610 | 49.3955 | 9.879 |
| 82 n-Butylbenzene | 91 | Compound Not Detected. | | | | | |
| 83 1,2-Dichlorobenzene | 146 | 10.295 | 10.295 | (1.039) | 1271863 | 47.8511 | 9.570 |
| 84 1,2-Dibromo-3-chloropropane | 157 | 11.052 | 11.064 | (1.116) | 129232 | 51.6742 | 10.335 |
| 85 1,2,4-Trichlorobenzene | 180 | 11.892 | 11.892 | (1.201) | 539429 | 47.6703 | 9.534 |
| 86 Hexachlorobutadiene | 225 | Compound Not Detected. | | | | | |
| 87 Naphthalene | 128 | Compound Not Detected. | | | | | |
| 88 1,2,3-Trichlorobenzene | 180 | Compound Not Detected. | | | | | |
| 98 Cyclohexane | 56 | 4.568 | 4.568 | (0.908) | 791178 | 45.7968 | 9.159 |
| 143 Methyl Acetate | 43 | 2.994 | 2.994 | (0.595) | 449301 | 46.5123 | 9.302 |
| 144 Methylcyclohexane | 83 | 5.514 | 5.514 | (1.096) | 649471 | 44.7998 | 8.960 |
| 141 1,3,5-Trichlorobenzene | 180 | Compound Not Detected. | | | | | |

Report Date: 04-Oct-2004 09:52

Calibration History

Method : \\qcanoh04\dd\chem\MSV\abux11.i\J41001A.b\8260LLUX11.m
Start Cal Date: 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Last Cal Level: 1
Last Cal Type : Initial Calibration

Initial Calibration

| Injection Date | Sublist | Calibration File |
|---|---------|------------------|
| Cal Level: 1 , Cal Amount: 5.000 | | |
| 14-SEP-2004 15:41 2-8260 UXJ23875.D | | |
| 16-AUG-2004 18:11 | 3-IX | UXJ23214.D |
| Cal Level: 2 , Cal Amount: 10.000 | | |
| 14-SEP-2004 15:19 2-8260 UXJ23874.D | | |
| 16-AUG-2004 17:48 | 3-IX | UXJ23213.D |
| Cal Level: 3 , Cal Amount: 25.000 | | |
| 14-SEP-2004 14:57 2-8260 UXJ23873.D | | |
| 16-AUG-2004 17:26 | 3-IX | UXJ23212.D |
| Cal Level: 4 , Cal Amount: 50.000 | | |
| 14-SEP-2004 14:33 2-8260 UXJ23872.D | | |
| 16-AUG-2004 17:03 | 3-IX | UXJ23211.D |
| Cal Level: 5 , Cal Amount: 100.00 | | |
| 14-SEP-2004 14:10 2-8260 UXJ23871.D | | |
| 16-AUG-2004 16:40 | 3-IX | UXJ23210.D |
| Cal Level: 6 , Cal Amount: 200.00 | | |
| 14-SEP-2004 13:48 2-8260 UXJ23870.D | | |
| 16-AUG-2004 16:18 | 3-IX | UXJ23209.D |

Continuing Calibration

| | | |
|-------------------|--------|------------|
| 01-OCT-2004 08:01 | 2-8260 | UXJ24277.D |
| 01-OCT-2004 08:24 | 3-IX | UXJ24278.D |

Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24277.D
 Report Date: 01-Oct-2004 08:48

STL North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: a3ux11.i Injection Date: 01-OCT-2004 08:01
 Lab File ID: UXJ24277.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
 Analysis Type: WATER Init. Cal. Times: 16:18 15:41
 Lab Sample ID: 50NG-CC Quant Type: ISTD
 Method: \\QCANOH04\dd\chem\MSV\ a3ux11.i\J41001A.b\8260LLUX11.m

| COMPOUND | RRF | RF50 | RRF | MIN | MAX |
|---------------------------------|----------|----------|-------|-------|------|
| | | | | %D | %D |
| 4 Dibromofluoromethane | 0.23279 | 0.25189 | 0.010 | 8.2 | 50.0 |
| 5 1,2-Dichloroethane-d4 | 0.32011 | 0.35221 | 0.010 | 10.0 | 50.0 |
| 6 Toluene-d8 | 1.20120 | 1.25338 | 0.010 | 4.3 | 50.0 |
| 7 Bromofluorobenzene | 0.51038 | 0.56722 | 0.010 | 11.1 | 50.0 |
| 8 Dichlorodifluoromethane | 0.26140 | 0.37793 | 0.010 | 44.6 | 50.0 |
| 9 Chloromethane | 0.46672 | 0.51868 | 0.100 | 11.1 | 50.0 |
| 10 Vinyl Chloride | 0.31925 | 0.35620 | 0.010 | 11.6 | 20.0 |
| 11 Bromomethane | 0.15143 | 0.11669 | 0.010 | -22.9 | 50.0 |
| 12 Chloroethane | 0.23463 | 0.26550 | 0.010 | 21.7 | 50.0 |
| 13 Trichlorofluoromethane | 0.33452 | 0.30267 | 0.010 | -9.5 | 50.0 |
| 15 Acrolein | 0.03123 | 0.04782 | 0.010 | 53.1 | 50.0 |
| 16 Acetone | 100 | 96.48724 | 0.010 | 3.5 | 50.0 |
| 17 1,1-Dichloroethene | 0.22360 | 0.23564 | 0.010 | 5.4 | 20.0 |
| 18 Freon-113 | 50.00000 | 55.87506 | 0.010 | -11.8 | 50.0 |
| 19 Iodomethane | 0.33050 | 0.25699 | 0.010 | -22.2 | 50.0 |
| 20 Carbon Disulfide | 0.83825 | 0.96631 | 0.010 | 15.7 | 50.0 |
| 21 Methylene Chloride | 50.00000 | 64.96049 | 0.010 | -29.9 | 50.0 |
| 22 Acetonitrile | 0.02967 | 0.04480 | 0.010 | 48.3 | 50.0 |
| 23 Acrylonitrile | 0.09228 | 0.11898 | 0.010 | -28.9 | 50.0 |
| 24 Methyl tert-butyl ether | 0.71686 | 0.67533 | 0.010 | -5.8 | 50.0 |
| 25 trans-1,2-Dichloroethene | 0.26526 | 0.27247 | 0.010 | 2.7 | 50.0 |
| 26 Hexane | 80.00000 | 47.91832 | 0.010 | 4.2 | 20.0 |
| 27 Vinyl acetate | 0.42446 | 0.50965 | 0.010 | 20.1 | 50.0 |
| 28 1,1-Dichloroethane | 0.48389 | 0.51298 | 0.100 | 6.0 | 50.0 |
| 29 tert-Butyl Alcohol | 0.01945 | 0.02077 | 0.010 | 6.8 | 50.0 |
| 30 2-Butanone | 0.13695 | 0.13589 | 0.010 | -0.8 | 50.0 |
| M 31 1,2-Dichloroethene (total) | 0.27198 | 0.20114 | 0.010 | 3.4 | 50.0 |
| 32 cis-1,2-dichloroethene | 0.27870 | 0.28981 | 0.010 | 4.0 | 50.0 |
| 33 2,2-Dichloropropane | 0.29161 | 0.17056 | 0.010 | -38.8 | 50.0 |
| 34 Bromochloromethane | 0.12906 | 0.14135 | 0.010 | 9.5 | 50.0 |
| 35 Chloroform | 0.48643 | 0.50499 | 0.010 | 3.8 | 20.0 |
| 36 Tetrahydrofuran | 50.00000 | 52.70191 | 0.010 | -5.4 | 50.0 |
| 37 1,1,1-Trichloroethane | 0.36737 | 0.28574 | 0.010 | -22.2 | 50.0 |
| 38 1,1-Dichloropropene | 0.34240 | 0.33286 | 0.010 | -2.8 | 50.0 |
| 39 Carbon Tetrachloride | 0.28845 | 0.26013 | 0.010 | -9.8 | 50.0 |
| 40 1,2-Dichloroethane | 0.39745 | 0.41834 | 0.010 | 5.3 | 50.0 |

Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24277.D
Report Date: 01-Oct-2004 08:48

STL North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: a3ux11.i Injection Date: 01-OCT-2004 08:01
Lab File ID: UXJ24277.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\ a3ux11.i\J41001A.b\8260LLUX11.m

| COMPOUND | RRF | RF50 | MIN | MAX |
|------------------------------|---------|---------------|-------------|-----|
| | | RRF | %D | %D |
| 41 Benzene | 1.15116 | 1.22204 0.010 | 6.2 50.0 | |
| 42 Trichloroethane | 0.26272 | 0.25830 0.010 | -1.7 50.0 | |
| 43 1,2-Dichloropropane | 0.28397 | 0.29058 0.010 | 2.3 20.0 | |
| 44 1,4-Dioxane | 0.00243 | 0.00311 0.010 | 27.9 50.0 | <- |
| 45 Dibromomethane | 0.15736 | 0.17455 0.010 | 10.9 50.0 | |
| 46 Bromodichloromethane | 0.37355 | 0.38290 0.010 | 2.5 50.0 | |
| 47 2-Chloroethyl vinyl ether | 0.15967 | 0.16468 0.010 | 3.1 50.0 | |
| 48 cis-1,3-Dichloropropene | 0.45706 | 0.43511 0.010 | -4.8 50.0 | |
| 49 4-Methyl-2-pentanone | 0.24577 | 0.32236 0.010 | 31.2 50.0 | |
| 50 Toluene | 1.49327 | 1.53382 0.010 | 2.7 20.0 | |
| 51 trans-1,3-Dichloropropene | 0.53539 | 0.49193 0.010 | -8.1 50.0 | |
| 52 Ethyl Methacrylate | 0.46066 | 0.48973 0.010 | 6.3 50.0 | |
| 53 1,1,2-Trichloroethane | 0.30290 | 0.33023 0.010 | 9.0 50.0 | |
| 54 1,3-Dichloropropane | 0.57071 | 0.59918 0.010 | 5.0 50.0 | |
| 55 Tetrachloroethene | 0.24193 | 0.24272 0.010 | 0.3 50.0 | |
| 56 2-Hexanone | 0.24179 | 0.29357 0.010 | 21.4 50.0 | |
| 57 dibromochloromethane | 0.32307 | 0.34691 0.010 | 7.4 50.0 | |
| 58 1,2-Dibromoethane | 0.29844 | 0.32456 0.010 | 8.8 50.0 | |
| 59 Chlorobenzene | 0.96688 | 1.00723 0.300 | 4.2 50.0 | |
| 60 1,1,1,2-Tetrachloroethane | 0.33590 | 0.34566 0.010 | 2.9 50.0 | |
| 61 Ethylbenzene | 0.48827 | 0.50019 0.010 | 2.4 20.0 | |
| 62 m + p-Xylene | 0.62888 | 0.65285 0.010 | 8.6 50.0 | |
| M 63 Xylenes (total) | 0.62505 | 0.67568 0.010 | -8.1 50.0 | |
| 64 Xylene-o | 0.61738 | 0.66133 0.010 | 7.1 50.0 | |
| 65 Styrene | 1.10404 | 1.25905 0.010 | 14.0 50.0 | |
| 66 Bromoform | 0.21960 | 0.25989 0.100 | 28.4 50.0 | |
| 67 Isopropylbenzene | 1.37503 | 1.42512 0.010 | 3.6 50.0 | |
| 68 1,1,2,2-Tetrachloroethane | 0.75371 | 0.83168 0.300 | 10.3 50.0 | |
| 69 1,4-Dichloro-2-butene | 0.24799 | 0.25606 0.010 | 3.3 50.0 | |
| 70 1,2,3-Trichloropropane | 0.24733 | 0.27349 0.010 | 10.6 50.0 | |
| 71 Bromobenzene | 0.74599 | 0.73556 0.010 | -1.4 50.0 | |
| 72 n-Propylbenzene | 0.69551 | 0.70005 0.010 | 0.7 50.0 | |
| 73 2-Chlorotoluene | 0.68879 | 0.69411 0.010 | 0.8 50.0 | |
| 74 1,3,5-Trimethylbenzene | 2.31439 | 2.31125 0.010 | -0.1 50.0 | |
| 75 4-Chlorotoluene | 0.73560 | 0.75720 0.010 | 2.9 50.0 | |
| 76 tert-Butylbenzene | 1.87499 | 1.67448 0.010 | -10.7 50.0 | |

Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J41001A.b\UXJ24277.D
Report Date: 01-Oct-2004 08:48

STL North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: A3UX11.i Injection Date: 01-OCT-2004 08:01
Lab File ID: UXJ24277.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\A3UX11.i\J41001A.b\8260LLUX11.m

| COMPOUND | RRF | MIN | | MAX | |
|--------------------------------|----------|----------|-------|-------|------|
| | | RF50 | RRF | tD | tD |
| 77 1,2,4-Trimethylbenzene | 2.47063 | 2.47134 | 0.010 | 0.0 | 50.0 |
| 78 sec-Butylbenzene | 2.53792 | 2.27952 | 0.010 | -10.2 | 50.0 |
| 79 4-Isopropyltoluene | 2.11151 | 1.95087 | 0.010 | -7.6 | 50.0 |
| 80 1,3-Dichlorobenzene | 1.37723 | 1.34984 | 0.010 | -2.0 | 50.0 |
| 81 1,4-Dichlorobenzene | 1.44089 | 1.44232 | 0.010 | 0.1 | 50.0 |
| 82 n-Butylbenzene | 1.88626 | 1.89367 | 0.010 | -15.5 | 50.0 |
| 83 1,2-Dichlorobenzene | 1.35648 | 1.33167 | 0.010 | -1.8 | 50.0 |
| 84 1,2-Dibromo-3-chloropropane | 0.12763 | 0.10788 | 0.010 | -15.5 | 50.0 |
| 85 1,2,4-Trichlorobenzene | 0.57750 | 0.32310 | 0.010 | -44.1 | 50.0 |
| 86 Hexachlorobutadiene | 50.00000 | 37.88578 | 0.010 | 24.2 | 50.0 |
| 87 Naphthalene | 1.36300 | 0.59504 | 0.010 | -56.3 | 50.0 |
| 88 1,2,3-Trichlorobenzene | 0.41413 | 0.20855 | 0.010 | -49.6 | 50.0 |
| 98 Cyclohexane | 0.36784 | 0.34136 | 0.010 | -7.2 | 50.0 |
| 143 Methyl Acetate | 0.20568 | 0.22613 | 0.010 | 9.9 | 50.0 |
| 144 Methylcyclohexane | 0.30868 | 0.25378 | 0.010 | -17.8 | 50.0 |
| 141 1,3,5-Trichlorobenzene | 0.76231 | 0.58138 | 0.010 | -23.7 | 50.0 |

Data File: \\qcانوH04\\dd\\chem\\MSV\\a3ux11.i\\J41001A.b\\UXJ24277.D
Report Date: 10/01/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24277.D
Analysis Type: WATER

Injection Date: 01-OCT-2004 08:01
Lab Sample ID: 50NG-CC
Method File: \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\J41001A

| COMPOUND | EXPECTED CONC. | MEASURED CONC. | %D | MAX % |
|------------------------------|-------------------|-------------------|------|----------|
| 0 Chlorobenzene | 50.0000 | 52.0870 | 4.2 | 50.0 |
| 0 Bromodichloromethane | 50.0000 | 51.2503 | 2.5 | 50.0 |
| 0 1,1,2,2-Tetrachloroethane | 50.0000 | 55.1722 | 10.3 | 50.0 |
| 0 Bromoform | 50.0000 | 59.1750 | 18.4 | 50.0 |
| 0 Styrene | 50.0000 | 57.0203 | 14.0 | 50.0 |
| 0 Xylene-o | 50.0000 | 53.5589 | 7.1 | 50.0 |
| 0 Xylenes (total) | 150.0000 | 162.1411 | 8.1 | 50.0 |
| 0 2-Hexanone | 100.0000 | 121.4129 | 21.4 | 50.0 |
| 0 Chloromethane | 50.0000 | 55.5671 | 11.1 | 50.0 |
| 0 Vinyl Chloride | 50.0000 | 55.7871 | 11.6 | 20.0 |
| 0 Bromomethane | 50.0000 | 38.5298 | 22.9 | 50.0 |
| 0 Chloroethane | 50.0000 | 60.8415 | 21.7 | 50.0 |
| 0 1,1-Dichloroethane | 50.0000 | 53.0059 | 6.0 | 50.0 |
| 0 Tetrachloroethene | 50.0000 | 50.1636 | 0.3 | 50.0 |
| 0 Acetone | 100.0000 | 96.4872 | 3.5 | 50.0 |
| 0 1,1-Dichloroethane | 50.0000 | 52.6915 | 5.4 | 20.0 |
| 0 m + p-Xylene | 100.0000 | 108.5822 | 8.6 | 50.0 |
| 0 Ethylbenzene | 50.0000 | 51.2208 | 2.4 | 20.0 |
| 0 Carbon Disulfide | 50.0000 | 57.8460 | 15.7 | 50.0 |
| 0 Methylene Chloride | 50.0000 | 64.9605 | 29.9 | 50.0 |
| 0 1,2-Dichloropropane | 50.0000 | 51.1639 | 2.3 | 20.0 |
| 0 1,1,2-Trichloroethane | 50.0000 | 54.5122 | 9.0 | 50.0 |
| 0 Dibromochloromethane | 50.0000 | 53.6891 | 7.4 | 50.0 |
| 0 trans-1,2-Dichloroethene | 50.0000 | 51.3593 | 2.7 | 50.0 |
| 0 trans-1,3-Dichloropropene | 50.0000 | 45.9410 | 8.1 | 50.0 |
| 0 cis-1,3-Dichloropropene | 50.0000 | 47.5991 | 4.8 | 50.0 |
| 0 Chloroform | 50.0000 | 51.9081 | 3.8 | 20.0 |
| 0 Toluene | 50.0000 | 51.3576 | 2.7 | 20.0 |
| 0 2-Butanone | 100.0000 | 99.2233 | 0.8 | 50.0 |
| 0 1,2-Dichloroethene (total) | 100.0000 | 103.3511 | 3.4 | 50.0 |
| 0 cis-1,2-dichloroethene | 50.0000 | 51.9918 | 4.0 | 50.0 |
| 0 4-Methyl-2-pentanone | 100.0000 | 131.1647 | 31.2 | 50.0 |
| 0 1,2-Dichloroethane | 50.0000 | 52.6284 | 5.3 | 50.0 |
| 0 Trichloroethene | 50.0000 | 49.1583 | 1.7 | 50.0 |
| 0 1,1,1-Trichloroethane | 50.0000 | 38.8901 | 22.2 | 50.0 |
| 0 Carbon Tetrachloride | 50.0000 | 45.0898 | 9.8 | 50.0 |
| 0 Benzene | 50.0000 | 53.0789 | 6.2 | 50.0 |
| 38 Dichlorodifluoromethane | 50.0000 | 72.2896 | 44.6 | 50.0 |
| 39 Trichlorofluoromethane | 50.0000 | 45.2263 | 9.5 | 50.0 |

Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b/UXJ24277.D
Report Date: 10/01/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24277.D
Analysis Type: WATER

Injection Date: 01-OCT-2004 08:01
Lab Sample ID: 50NG-CC
Method File: \\QCANOH04\dd\chem\MSV\ a3ux11.i\

| COMPOUND | EXPECTED CONC. | MEASURED CONC. | %D | MAX % |
|--------------------------------|-------------------|-------------------|------|----------|
| 39 Chlorobenzene-d5 | 50.0000 | 50.0000 | 0.0 | 50.0 |
| 40 Acrolein | 500.0000 | 765.5361 | 53.1 | 50.0 <- |
| 41 Acrylonitrile | 500.0000 | 644.6819 | 28.9 | 50.0 |
| 42 Vinyl acetate | 50.0000 | 60.0356 | 20.1 | 50.0 |
| 43 2-Chloroethyl vinyl ether | 100.0000 | 103.1382 | 3.1 | 50.0 |
| 47 Freon-113 | 50.0000 | 55.8751 | 11.8 | 50.0 |
| 48 1,3-Dichlorobenzene | 50.0000 | 49.0056 | 2.0 | 50.0 |
| 49 1,4-Dichlorobenzene | 50.0000 | 50.0497 | 0.1 | 50.0 |
| 50 1,2-Dichlorobenzene | 50.0000 | 49.0857 | 1.8 | 50.0 |
| 51 Acetonitrile | 500.0000 | 741.3811 | 48.3 | 50.0 |
| 52 Iodomethane | 50.0000 | 38.8784 | 22.2 | 50.0 |
| 59 1,4-Dioxane | 2500.0000 | 3197.8748 | 27.9 | 50.0 |
| 60 Dibromomethane | 50.0000 | 55.4632 | 10.9 | 50.0 |
| 62 Ethyl Methacrylate | 50.0000 | 53.1558 | 6.3 | 50.0 |
| 63 1,2-Dibromoethane | 50.0000 | 54.3773 | 8.8 | 50.0 |
| 64 1,1,1,2-Tetrachloroethane | 50.0000 | 51.4523 | 2.9 | 50.0 |
| 65 1,2,3-Trichloropropane | 50.0000 | 55.2876 | 10.6 | 50.0 |
| 66 1,4-Dichloro-2-butene | 50.0000 | 51.6264 | 3.3 | 50.0 |
| 69 1,2-Dibromo-3-chloropropane | 50.0000 | 42.2635 | 15.6 | 50.0 |
| 82 Methyl tert-butyl ether | 50.0000 | 47.1031 | 5.8 | 50.0 |
| 84 Tetrahydrofuran | 50.0000 | 52.7019 | 5.4 | 50.0 |
| 98 2,2-Dichloropropane | 50.0000 | 30.6172 | 38.8 | 50.0 |
| 99 1,1-Dichloropropene | 50.0000 | 48.6079 | 2.8 | 50.0 |
| 100 1,3-Dichloropropane | 50.0000 | 52.4938 | 5.0 | 50.0 |
| 102 Bromobenzene | 50.0000 | 49.3006 | 1.4 | 50.0 |
| 103 2-Chlorotoluene | 50.0000 | 50.3866 | 0.8 | 50.0 |
| 104 n-Propylbenzene | 50.0000 | 50.3260 | 0.7 | 50.0 |
| 105 4-Chlorotoluene | 50.0000 | 51.4545 | 2.9 | 50.0 |
| 106 1,3,5-Trimethylbenzene | 50.0000 | 49.9323 | 0.1 | 50.0 |
| 107 tert-Butylbenzene | 50.0000 | 44.6530 | 10.7 | 50.0 |
| 108 1,2,4-Trimethylbenzene | 50.0000 | 50.0142 | 0.0 | 50.0 |
| 109 sec-Butylbenzene | 50.0000 | 44.9052 | 10.2 | 50.0 |
| 110 4-Isopropyltoluane | 50.0000 | 46.1961 | 7.6 | 50.0 |
| 111 n-Butylbenzene | 50.0000 | 42.2440 | 15.5 | 50.0 |
| 112 1,2,4-Trichlorobenzene | 50.0000 | 27.9738 | 44.1 | 50.0 |
| 113 Naphthalene | 50.0000 | 21.8283 | 56.3 | 50.0 <- |
| 114 Hexachlorobutadiene | 50.0000 | 37.8858 | 24.2 | 50.0 |
| 115 1,2,3-Trichlorobenzene | 50.0000 | 25.1796 | 49.6 | 50.0 |
| 124 tert-Butyl Alcohol | 1000.0000 | 1067.9313 | 6.8 | 50.0 |

Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J41001A.b/UXJ24277.D
Report Date: 10/01/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24277.D
Analysis Type: WATER

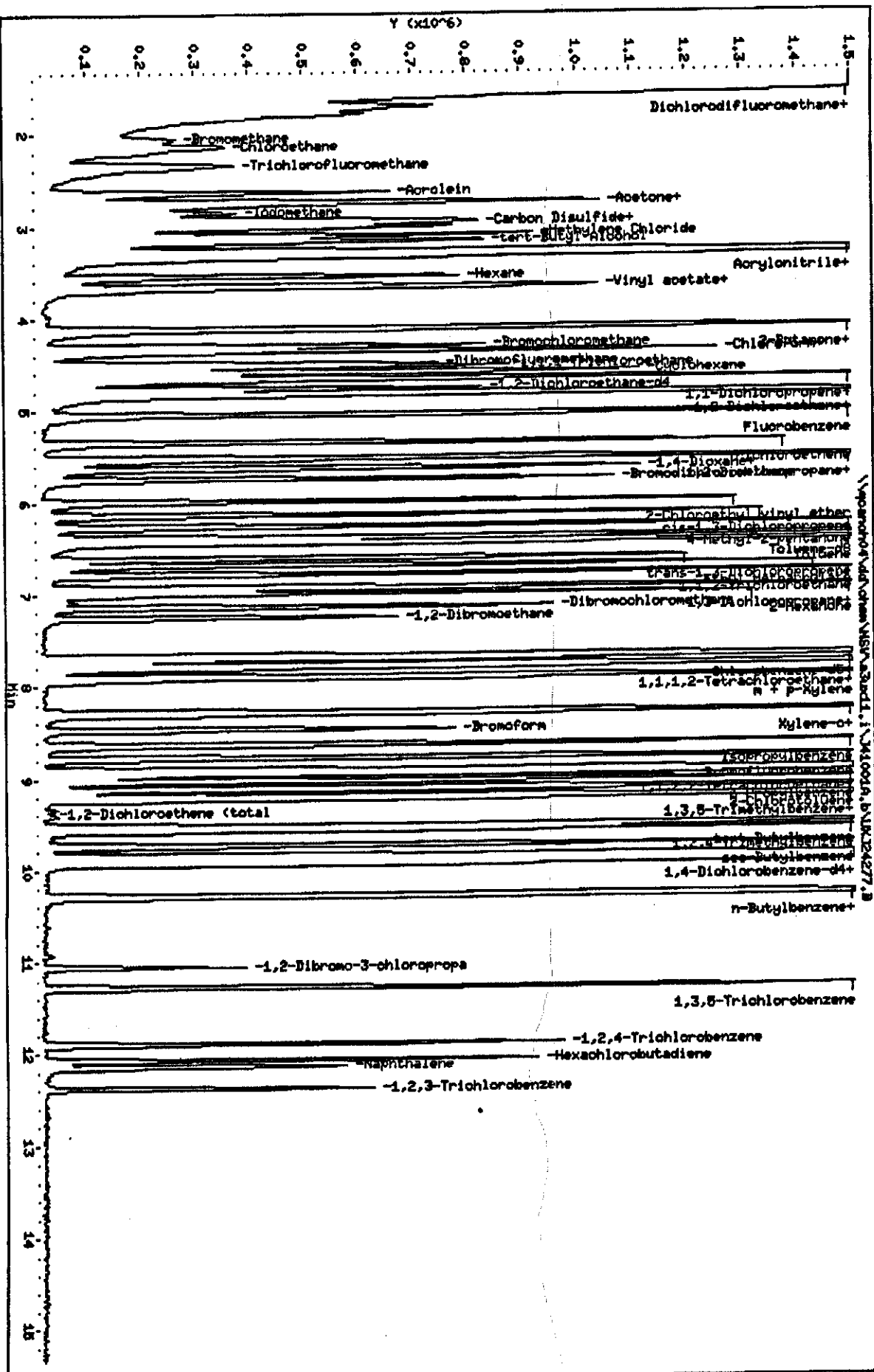
Injection Date: 01-OCT-2004 08:01
Lab Sample ID: 50NG-CC
Method File: \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\

| COMPOUND | EXPECTED | MEASURED | %D | MAX |
|----------------------------|----------|----------|------|------|
| | CONC. | CONC. | | |
| 125 Hexane | 50.0000 | 47.9183 | 4.3 | 20.0 |
| 127 Cyclohexane | 50.0000 | 46.4012 | 7.2 | 50.0 |
| 128 Isopropylbenzene | 50.0000 | 51.8215 | 3.6 | 50.0 |
| 130 Fluorobenzene | 50.0000 | 50.0000 | 0.0 | 50.0 |
| 132 1,4-Dichlorobenzene-d4 | 50.0000 | 50.0000 | 0.0 | 50.0 |
| 133 Bromochloromethane | 50.0000 | 54.7609 | 9.5 | 50.0 |
| 141 1,3,5-Trichlorobenzene | 50.0000 | 38.1325 | 23.7 | 50.0 |
| 143 Methyl Acetate | 100.0000 | 109.9442 | 9.9 | 50.0 |
| 144 Methylcyclohexane | 50.0000 | 41.1082 | 17.8 | 50.0 |
| 22 Toluene-d8 | 50.0000 | 52.1721 | 4.3 | 50.0 |
| 32 Bromofluorobenzene | 50.0000 | 55.5686 | 11.1 | 50.0 |
| 47 1,2-Dichloroethane-d4 | 50.0000 | 55.0141 | 10.0 | 50.0 |
| 131 Dibromofluoromethane | 50.0000 | 54.1014 | 8.2 | 50.0 |

Date File: \\pcpanth04\dat\chem\NIST\as30d1.\J3410019.b\IRK24277.D
Date : 01-OCT-2004 08:01
Client ID:
Sample Info: SONG-CC
Purge Volume: 5.0
Column phase: IRD24

Instrument: as30d1.i

Operator: 43982
Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24277.D
Report Date: 04-Oct-2004 09:54

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24277.D
Lab Smp Id: 50NG-CC
Inj Date : 01-OCT-2004 08:01
Operator : 43582
Smp Info : 50NG-CC
Misc Info : J41001A,8260LLUX11,2-8260.SUB,43582,2
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 1 Continuing Calibration Sample
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|-----------|---------------------------|---------|-------|---------------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 2028310 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1659318 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 1038318 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 510911 | 50.0000 | 54.101 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 714395 | 50.0000 | 55.014 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 2079759 | 50.0000 | 52.172 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 941200 | 50.0000 | 55.569 | |
| \$ | 8 Dichlorodifluoromethane | 85 | 1.550 | 1.550 (0.308) | 766557 | 50.0000 | 72.290 | |
| 9 | Chloromethane | 50 | 1.704 | 1.704 (0.338) | 1052052 | 50.0000 | 55.567 | |
| 10 | Vinyl Chloride | 62 | 1.787 | 1.787 (0.355) | 722492 | 50.0000 | 55.787 | |
| 11 | Bromomethane | 94 | 2.059 | 2.059 (0.409) | 236682 | 50.0000 | 38.530 | |
| 12 | Chloroethane | 64 | 2.142 | 2.142 (0.425) | 579087 | 50.0000 | 60.842 | |
| 13 | Trichlorofluoromethane | 101 | 2.331 | 2.331 (0.463) | 613915 | 50.0000 | 45.226 | |
| 15 | Acrolein | 56 | 2.639 | 2.639 (0.524) | 969923 | 500.000 | 765.54 | |
| 16 | Acetone | 43 | 2.745 | 2.745 (0.545) | 437054 | 100.000 | 96.487 | |
| 17 | 1,1-Dichloroethane | 96 | 2.734 | 2.734 (0.542) | 477951 | 50.0000 | 52.692 | |
| 18 | Freon-113 | 151 | 2.745 | 2.745 (0.545) | 318328 | 50.0000 | 55.875 | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24277.D
 Report Date: 04-Oct-2004 09:54

| Compounds | QUANT SIG | AMOUNTS | | | | | |
|---------------------------------|-----------|---------|---------------|---------|---------|------------|------------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) |
| 19 Iodomethane | 142 | 2.864 | 2.864 (0.568) | 521247 | 50.0000 | 38.878 | |
| 20 Carbon Disulfide | 76 | 2.923 | 2.923 (0.580) | 1959986 | 50.0000 | 57.846 | |
| 21 Methylene Chloride | 84 | 3.100 | 3.100 (0.615) | 801851 | 50.0000 | 64.960 | |
| 22 Acetonitrile | 41 | 2.958 | 2.958 (0.587) | 892476 | 500.000 | 741.38 | |
| 23 Acrylonitrile | 53 | 3.278 | 3.278 (0.650) | 2413328 | 500.000 | 644.68 | |
| 24 Methyl tert-butyl ether | 73 | 3.325 | 3.325 (0.660) | 1369770 | 50.0000 | 47.103 | |
| 25 trans-1,2-Dichloroethane | 96 | 3.325 | 3.325 (0.660) | 552652 | 50.0000 | 51.359 | |
| 26 Hexane | 86 | 3.550 | 3.550 (0.704) | 83387 | 50.0000 | 47.918 | |
| 27 Vinyl acetate | 43 | 3.680 | 3.680 (0.730) | 1033735 | 50.0000 | 60.036 | |
| 28 1,1-Dichloroethane | 63 | 3.657 | 3.657 (0.725) | 1040482 | 50.0000 | 53.006 | |
| 29 tert-Butyl Alcohol | 59 | 3.171 | 3.171 (0.629) | 842515 | 1000.00 | 1067.9 (A) | |
| 30 2-Butanone | 43 | 4.106 | 4.106 (0.815) | 551238 | 100.000 | 99.223 | |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 1140469 | 100.000 | 103.35 |
| 32 cis-1,2-dichloroethene | 96 | 4.106 | 4.106 (0.815) | 587817 | 50.0000 | 51.992 | |
| 33 2,2-Dichloropropane | 77 | 4.118 | 4.118 (0.817) | 362182 | 50.0000 | 30.617 | |
| 34 Bromochloromethane | 128 | 4.296 | 4.296 (0.852) | 286692 | 50.0000 | 54.761 | |
| 35 Chloroform | 83 | 4.355 | 4.355 (0.864) | 1024286 | 50.0000 | 51.908 | |
| 36 Tetrahydrofuran | 42 | 4.343 | 4.343 (0.862) | 161153 | 50.0000 | 52.702 | |
| 37 1,1,1-Trichloroethane | 97 | 4.520 | 4.520 (0.897) | 579574 | 50.0000 | 38.890 | |
| 38 1,1-Dichloropropene | 75 | 4.651 | 4.651 (0.923) | 675150 | 50.0000 | 48.608 | |
| 39 Carbon Tetrachloride | 117 | 4.662 | 4.662 (0.925) | 527617 | 50.0000 | 45.090 | |
| 40 1,2-Dichloroethane | 62 | 4.816 | 4.816 (0.955) | 848519 | 50.0000 | 52.628 | |
| 41 Benzene | 78 | 4.828 | 4.828 (0.958) | 2478681 | 50.0000 | 53.079 | |
| 42 Trichloroethene | 130 | 5.349 | 5.349 (1.061) | 523914 | 50.0000 | 49.158 | |
| 43 1,2-Dichloropropane | 63 | 5.526 | 5.526 (1.096) | 589385 | 50.0000 | 51.164 | |
| 44 1,4-Dioxane | 88 | 5.633 | 5.633 (1.117) | 315764 | 2500.00 | 3197.9 (A) | |
| 45 Dibromomethane | 93 | 5.621 | 5.621 (1.118) | 354039 | 50.0000 | 55.463 | |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 (1.141) | 776631 | 50.0000 | 51.250 | |
| 47 2-Chloroethyl vinyl ether | 63 | 5.999 | 5.999 (1.190) | 668047 | 100.000 | 103.14 | |
| 48 cis-1,3-Dichloropropene | 75 | 6.130 | 6.130 (1.216) | 882542 | 50.0000 | 47.599 | |
| 49 4-Methyl-2-pentanone | 43 | 6.248 | 6.248 (1.239) | 1307685 | 100.000 | 131.16 | |
| 50 Toluene | 91 | 6.437 | 6.437 (0.838) | 2545089 | 50.0000 | 51.358 | |
| 51 trans-1,3-Dichloropropene | 75 | 6.615 | 6.615 (0.861) | 816264 | 50.0000 | 45.941 | |
| 52 Ethyl Methacrylate | 69 | 6.686 | 6.686 (0.871) | 812621 | 50.0000 | 53.156 | |
| 53 1,1,2-Trichloroethane | 97 | 6.780 | 6.780 (0.883) | 547357 | 50.0000 | 54.512 | |
| 54 1,3-Dichloropropane | 76 | 6.922 | 6.922 (0.901) | 994224 | 50.0000 | 52.494 | |
| 55 Tetrachloroethane | 164 | 6.934 | 6.934 (0.903) | 402750 | 50.0000 | 50.164 | |
| 56 2-Hexanone | 43 | 6.993 | 6.993 (0.911) | 974244 | 100.000 | 121.41 | |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 (0.929) | 575628 | 50.0000 | 53.689 | |
| 58 1,2-Dibromoethane | 107 | 7.254 | 7.254 (0.945) | 538554 | 50.0000 | 54.377 | |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 (1.003) | 1671322 | 50.0000 | 52.087 | |
| 60 1,1,1,2-Tetrachloroethane | 131 | 7.774 | 7.774 (1.012) | 573562 | 50.0000 | 51.452 | |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 (1.015) | 829982 | 50.0000 | 51.221 | |
| 62 m + p-Xylene | 106 | 7.905 | 7.905 (1.029) | 2266141 | 100.000 | 108.58 | |
| M 63 Xylenes (total) | 106 | 8.283 | 8.283 (1.079) | 1097349 | 50.0000 | 53.559 | |
| 64 Xylene-o | 106 | 8.295 | 8.295 (1.080) | 2089170 | 50.0000 | 57.020 | |
| 65 Styrene | 104 | | | | | | |

Data File: \\qcanoh04\dd\chem\MSV\aux11.i\J41001A.b\UXJ24277.D
 Report Date: 04-Oct-2004 09:54

| Compounds | QUANT SIG | MASS | RT | EXP RT | REL RT | RESPONSE | AMOUNTS | |
|--------------------------------|-----------|------|--------|----------------|--------|----------|------------------|-----------------|
| | | | | | | | CAL-AMT (ng) | ON-COL (ng) |
| 66 Bromoform | | 173 | 8.473 | 8.473 (1.103) | | 431247 | 50.0000 | 59.175 |
| 67 Isopropylbenzene | | 105 | 8.626 | 8.626 (1.123) | | 2364729 | 50.0000 | 51.821 |
| 68 1,1,2,2-Tetrachloroethane | | 83 | 8.899 | 8.899 (0.898) | | 863546 | 50.0000 | 55.172 |
| 69 1,4-Dichloro-2-butene | | 53 | 8.958 | 8.958 (0.904) | | 265867 | 50.0000 | 51.626 |
| 70 1,2,3-Trichloropropane | | 110 | 8.946 | 8.946 (0.903) | | 283957 | 50.0000 | 55.288 |
| 71 Bromobenzene | | 156 | 8.934 | 8.934 (0.902) | | 763740 | 50.0000 | 49.300 |
| 72 n-Propylbenzene | | 120 | 9.029 | 9.029 (0.912) | | 726874 | 50.0000 | 50.326 |
| 73 2-Chlorotoluene | | 126 | 9.112 | 9.112 (0.920) | | 720708 | 50.0000 | 50.386 |
| 74 1,3,5-Trimethylbenzene | | 105 | 9.194 | 9.194 (0.928) | | 2399817 | 50.0000 | 49.932 |
| 75 4-Chlorotoluene | | 126 | 9.218 | 9.218 (0.931) | | 786217 | 50.0000 | 51.454 |
| 76 tert-Butylbenzene | | 119 | 9.514 | 9.514 (0.961) | | 1738644 | 50.0000 | 44.653 |
| 77 1,2,4-Trimethylbenzene | | 105 | 9.561 | 9.561 (0.965) | | 2566033 | 50.0000 | 50.014 |
| 78 sec-Butylbenzene | | 105 | 9.727 | 9.727 (0.982) | | 2366867 | 50.0000 | 44.909 |
| 79 4-Isopropyltoluene | | 119 | 9.869 | 9.869 (0.996) | | 2025619 | 50.0000 | 46.196 |
| 80 1,3-Dichlorobenzene | | 146 | 9.845 | 9.845 (0.994) | | 1401559 | 50.0000 | 49.006 |
| 81 1,4-Dichlorobenzene | | 146 | 9.928 | 9.928 (1.002) | | 1497588 | 50.0000 | 50.050 |
| 82 n-Butylbenzene | | 91 | 10.271 | 10.271 (1.037) | | 1654733 | 50.0000 | 42.244 |
| 83 1,2-Dichlorobenzene | | 146 | 10.295 | 10.295 (1.039) | | 1382698 | 50.0000 | 49.086 |
| 84 1,2-Dibromo-3-chloropropane | | 157 | 11.052 | 11.052 (1.116) | | 112018 | 50.0000 | 42.264 |
| 85 1,2,4-Trichlorobenzene | | 180 | 11.892 | 11.892 (1.201) | | 335477 | 50.0000 | 27.974 |
| 86 Hexachlorobutadiene | | 225 | 12.070 | 12.070 (1.219) | | 203779 | 50.0000 | 37.886 |
| 87 Naphthalene | | 128 | 12.129 | 12.129 (1.225) | | 617838 | 50.0000 | 21.828 |
| 88 1,2,3-Trichlorobenzene | | 180 | 12.377 | 12.377 (1.250) | | 216543 | 50.0000 | 25.180 |
| 98 Cyclohexane | | 56 | 4.580 | 4.580 (0.908) | | 692392 | 50.0000 | 46.401 |
| 143 Methyl Acetate | | 43 | 3.006 | 3.006 (0.596) | | 917329 | 100.000 | 109.94 |
| 144 Methylcyclohexane | | 83 | 5.526 | 5.526 (1.096) | | 514749 | 50.0000 | 41.108 |
| 141 1,3,5-Trichlorobenzene | | 180 | 11.277 | 11.277 (1.139) | | 603654 | 50.0000 | 38.132 |

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\QCANOH04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24278.D
Report Date: 01-Oct-2004 08:40

STL - North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: a3ux11.i Injection Date: 01-OCT-2004 08:24
Lab File ID: UXJ24278.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-A9CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\ a3ux11.i\J41001A.b\8260LLUX11.m

| COMPOUND | RRF | RF50 | MIN | MAX |
|---------------------------|---------|---------------|-------|---------|
| | | RRF | %D | %D |
| 14 Dichlorofluoromethane | 0.48495 | 0.65672 0.010 | 35.4 | 50.0 |
| 89 Ethyl Ether | 0.24654 | 0.24329 0.010 | -1.3 | 50.0 |
| 91 3-Chloropropene | 0.10305 | 0.13656 0.010 | 32.5 | 50.0 |
| 92 Isopropyl Ether | 0.22353 | 0.23717 0.010 | 6.1 | 50.0 |
| 93 2-Chloro-1,3-butadiene | 0.37276 | 0.42189 0.010 | 13.2 | 50.0 |
| 94 Propionitrile | 0.04231 | 0.04179 0.010 | -1.2 | 50.0 |
| 95 Ethyl Acetate | 0.24508 | 0.25242 0.010 | 3.0 | 50.0 |
| 96 Methacrylonitrile | 0.15890 | 0.16351 0.010 | 2.9 | 50.0 |
| 97 Isobutanol | 0.01142 | 0.01217 0.010 | 6.5 | 50.0 |
| 99 n-Butanol | 0.00822 | 0.00762 0.010 | -7.3 | 50.0 <- |
| 100 Methyl Methacrylate | 0.19531 | 0.21671 0.010 | 11.0 | 50.0 |
| 101 2-Nitropropane | 0.06079 | 0.06355 0.010 | 4.5 | 50.0 |
| 103 Cyclohexanone | 0.02717 | 0.02360 0.010 | -13.2 | 50.0 |

Data File: \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\J41001A.b\\UXJ24278.D
Report Date: 10/01/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24278.D
Analysis Type: WATER

Injection Date: 01-OCT-2004 08:24
Lab Sample ID: 50NG-A9CC
Method File: \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\J41001A

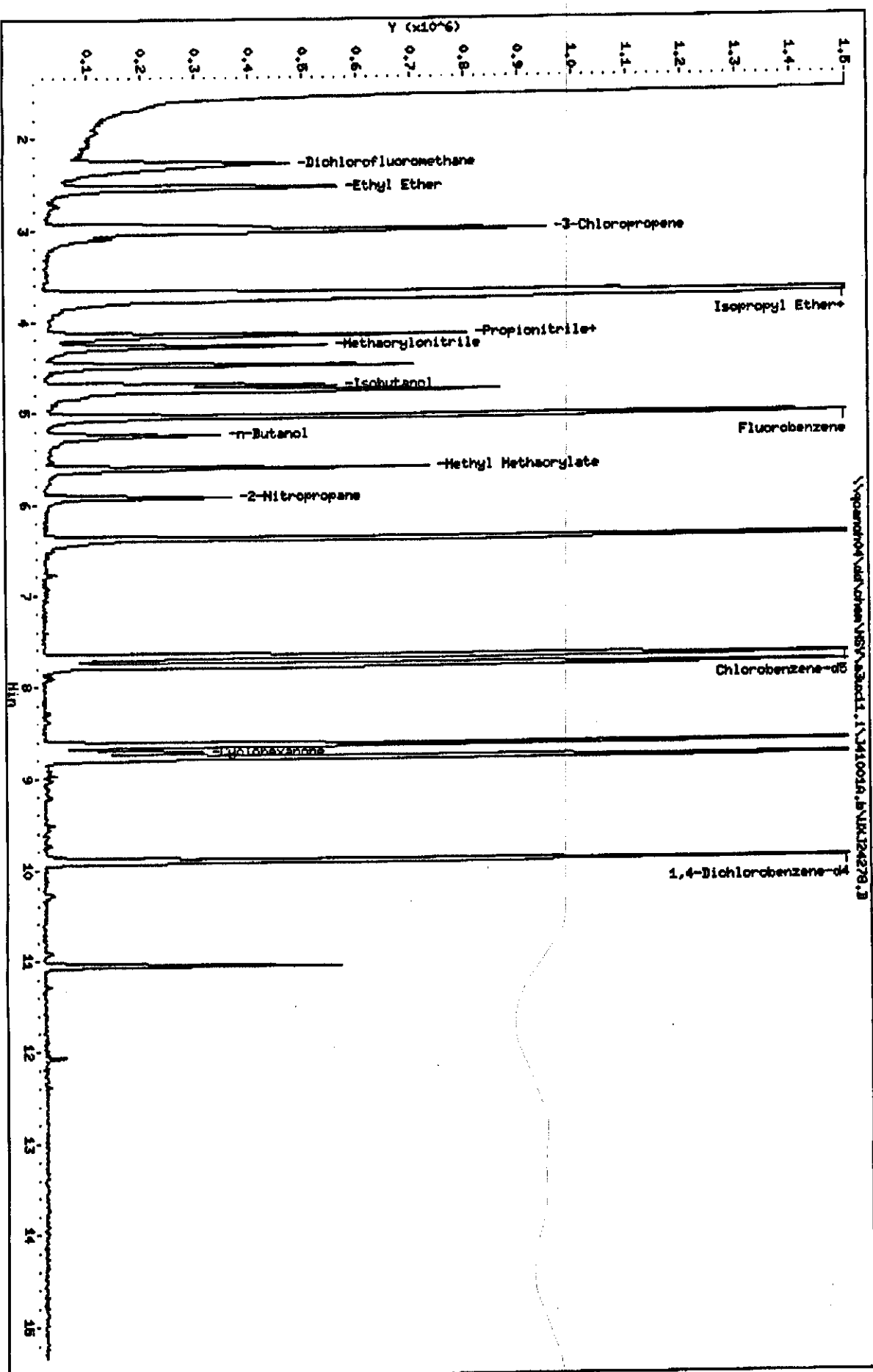
| COMPOUND | EXPECTED | MEASURED | %D | MAX |
|----------------------------|-----------|-----------|------|------|
| | CONC. | CONC. | | |
| 39 Chlorobenzene-d5 | 50.0000 | 50.0000 | 0.0 | 50.0 |
| 53 3-Chloropropene | 50.0000 | 66.2612 | 32.5 | 50.0 |
| 54 2-Chloro-1,3-butadiene | 50.0000 | 56.5904 | 13.2 | 50.0 |
| 55 Propionitrile | 100.0000 | 98.7794 | 1.2 | 50.0 |
| 56 Methacrylonitrile | 50.0000 | 51.4506 | 2.9 | 50.0 |
| 57 Isobutanol | 1000.0000 | 1065.4288 | 6.5 | 50.0 |
| 58 Methyl Methacrylate | 50.0000 | 55.4790 | 11.0 | 50.0 |
| 73 n-Butanol | 1000.0000 | 926.6878 | 7.3 | 50.0 |
| 74 Ethyl Acetate | 100.0000 | 102.9919 | 3.0 | 50.0 |
| 75 Cyclohexanone | 500.0000 | 434.1770 | 13.2 | 50.0 |
| 76 Ethyl Ether | 50.0000 | 49.3397 | 1.3 | 50.0 |
| 85 Dichlorofluoromethane | 50.0000 | 67.7105 | 35.4 | 50.0 |
| 86 2-Nitropropane | 100.0000 | 104.5366 | 4.5 | 50.0 |
| 126 Isopropyl Ether | 250.0000 | 265.2542 | 6.1 | 50.0 |
| 130 Fluorobenzene | 50.0000 | 50.0000 | 0.0 | 50.0 |
| 132 1,4-Dichlorobenzene-d4 | 50.0000 | 50.0000 | 0.0 | 50.0 |

Data File: \\pcando4\adachem\MSI\subdir.1\110019.d\\MSI24278.D
Date : 01-07-2004 08:24
Client ID:
Sample Info: SONG-A9CC
Purge Volume: 5.0
Column phase: 30624

Instrument: 30624.i

Operator: 43682
Column diameter: 0.18

\\pcando4\adachem\MSI\subdir.1\110019.d\\MSI24278.D



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24278.D
Report Date: 01-Oct-2004 08:47

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24278.D
Lab Smp Id: 50NG-A9CC
Inj Date : 01-OCT-2004 08:24
Operator : 43582
Smp Info : 50NG-A9CC
Misc Info : J41001A,8260LLUX11,3-IX.SUB,43582,2
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 01-Oct-2004 08:47 evans1 Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 2 Continuing Calibration Sample
Dil Factor: 1.00000
Integrator: HP RTE
Target Version: 4.04
Processing Host: CANPMSV07
Compound Sublist: 3-IX.SUB

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | AMOUNTS | | | | | | |
|-----------|---------------------------|---------|-------|---------------|---------|----------|------------------|-----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | CAL-AMT (ng) | ON-COL (ng) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 2032270 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1610209 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 736452 | 50.0000 | | |
| | 14 Dichlorofluoromethane | 67 | 2.284 | 2.284 (0.453) | 1334638 | 50.0000 | 67.710 | |
| | 89 Ethyl Ether | 59 | 2.532 | 2.532 (0.502) | 494425 | 50.0000 | 49.340 | |
| | 91 3-Chloropropene | 76 | 3.006 | 3.006 (0.596) | 277536 | 50.0000 | 66.261 | |
| | 92 Isopropyl Ether | 87 | 3.704 | 3.704 (0.735) | 2409975 | 250.000 | 265.25(A) | |
| | 93 2-Chloro-1,3-butadiene | 53 | 3.728 | 3.728 (0.739) | 857391 | 50.0000 | 56.590 | |
| | 94 Propionitrile | 54 | 4.142 | 4.142 (0.822) | 169864 | 100.000 | 98.779 | |
| | 95 Ethyl Acetate | 43 | 4.154 | 4.154 (0.824) | 1028952 | 100.000 | 102.99 | |
| | 96 Methacrylonitrile | 41 | 4.272 | 4.272 (0.847) | 332289 | 50.0000 | 51.450 | |
| | 97 Isobutanol | 41 | 4.710 | 4.710 (0.613) | 191774 | 1000.00 | 1065.4(A) | |
| | 99 n-Butanol | 56 | 5.254 | 5.254 (0.684) | 245256 | 1000.00 | 926.69(A) | |
| | 100 Methyl Methacrylate | 41 | 5.609 | 5.609 (1.113) | 440411 | 50.0000 | 55.479 | |
| | 101 2-Nitropropane | 41 | 5.928 | 5.928 (1.176) | 256312 | 100.000 | 104.54 | |
| | 103 Cyclohexanone | 55 | 8.709 | 8.709 (0.879) | 173768 | 500.000 | 434.17(A) | |

Data File: \\gcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24278.D
Report Date: 01-Oct-2004 08:47

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

SEVERN
TRENT

STL

RAW QC DATA

Date : 16-AUG-2004 13:09

Client ID: 50NG BFB

Instrument: z3ux11.i

Sample Info:

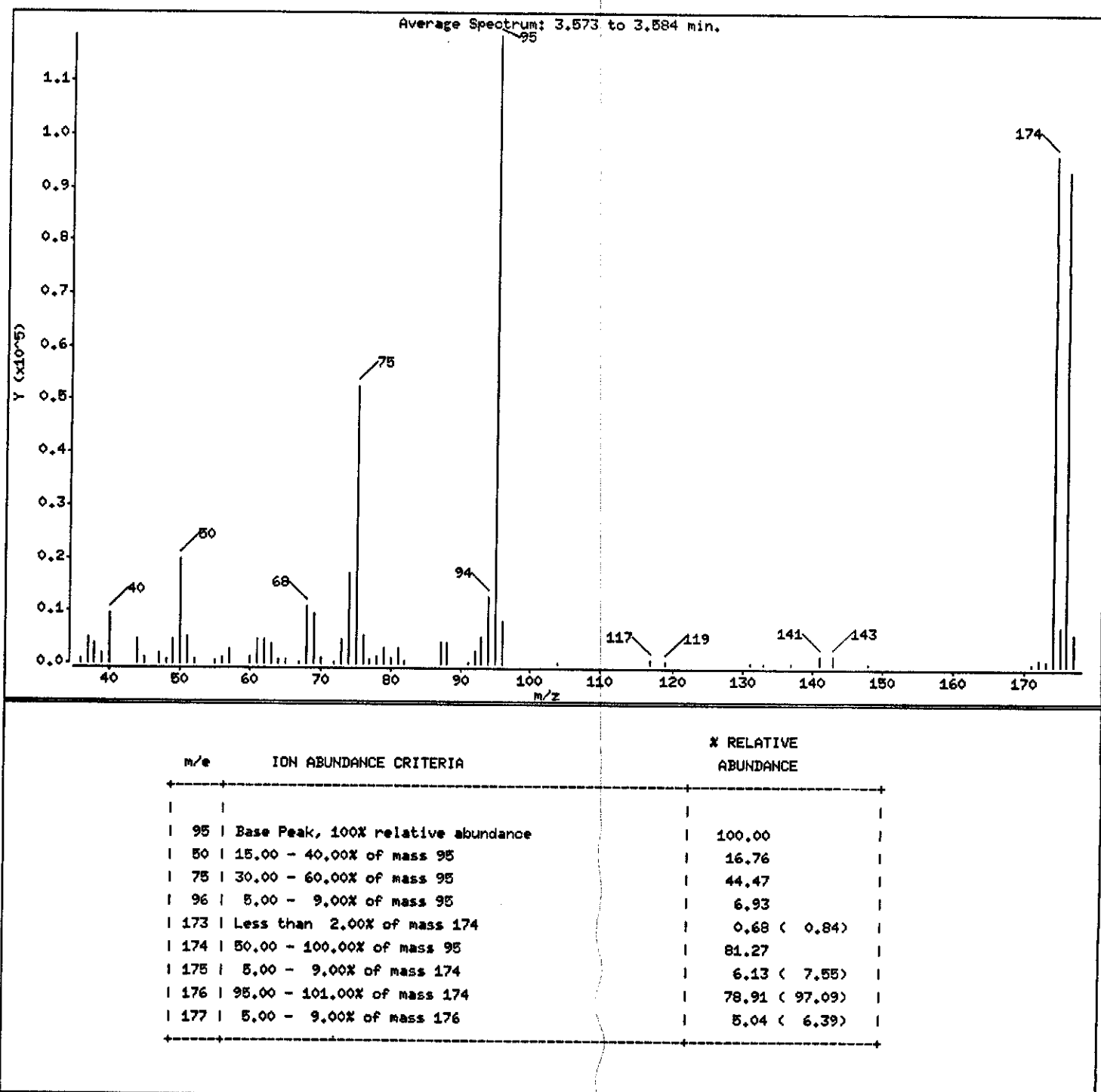
Volume Injected (uL): 1.0

Operator: 43882

Column phase: DB624 20M

Column diameter: 0.18

1 bfb



Date : 16-AUG-2004 13:09

Client ID: 5ONG BFB

Instrument: z3ux11.i

Sample Info:

Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

Data File: BFB207.D

Spectrum: Average Spectrum: 3.573 to 3.584 min.

Location of Maximum: 95.00

Number of points: 61

| m/z | Y | m/z | Y | m/z | Y | m/z | Y |
|-------|-------|-------|-------|--------|--------|--------|-------|
| 36.00 | 749 | 60.00 | 1376 | 78.00 | 1307 | 131.00 | 285 |
| 37.00 | 4867 | 61.00 | 4538 | 79.00 | 3095 | 133.00 | 271 |
| 38.00 | 3795 | 62.00 | 4718 | 80.00 | 1085 | 137.00 | 323 |
| 39.00 | 1854 | 63.00 | 3709 | 81.00 | 3055 | 141.00 | 1521 |
| 40.00 | 9546 | 64.00 | 700 | 82.00 | 515 | 143.00 | 1655 |
| 44.00 | 4571 | 65.00 | 821 | 87.00 | 4063 | 148.00 | 297 |
| 45.00 | 1190 | 67.00 | 254 | 88.00 | 4069 | 171.00 | 316 |
| 47.00 | 2040 | 68.00 | 10867 | 91.00 | 257 | 172.00 | 1062 |
| 48.00 | 737 | 69.00 | 9493 | 92.00 | 2338 | 173.00 | 806 |
| 49.00 | 4592 | 70.00 | 1139 | 93.00 | 5215 | 174.00 | 96288 |
| 50.00 | 19856 | 72.00 | 389 | 94.00 | 12726 | 175.00 | 7267 |
| 51.00 | 5302 | 73.00 | 4639 | 95.00 | 118480 | 176.00 | 93488 |
| 52.00 | 840 | 74.00 | 17288 | 96.00 | 8207 | 177.00 | 8974 |
| 55.00 | 613 | 75.00 | 52680 | 104.00 | 385 | | |
| 56.00 | 1116 | 76.00 | 5372 | 117.00 | 715 | | |
| 57.00 | 2804 | 77.00 | 828 | 119.00 | 414 | | |

Sample Info:

Volume Injected (uL): 1.0

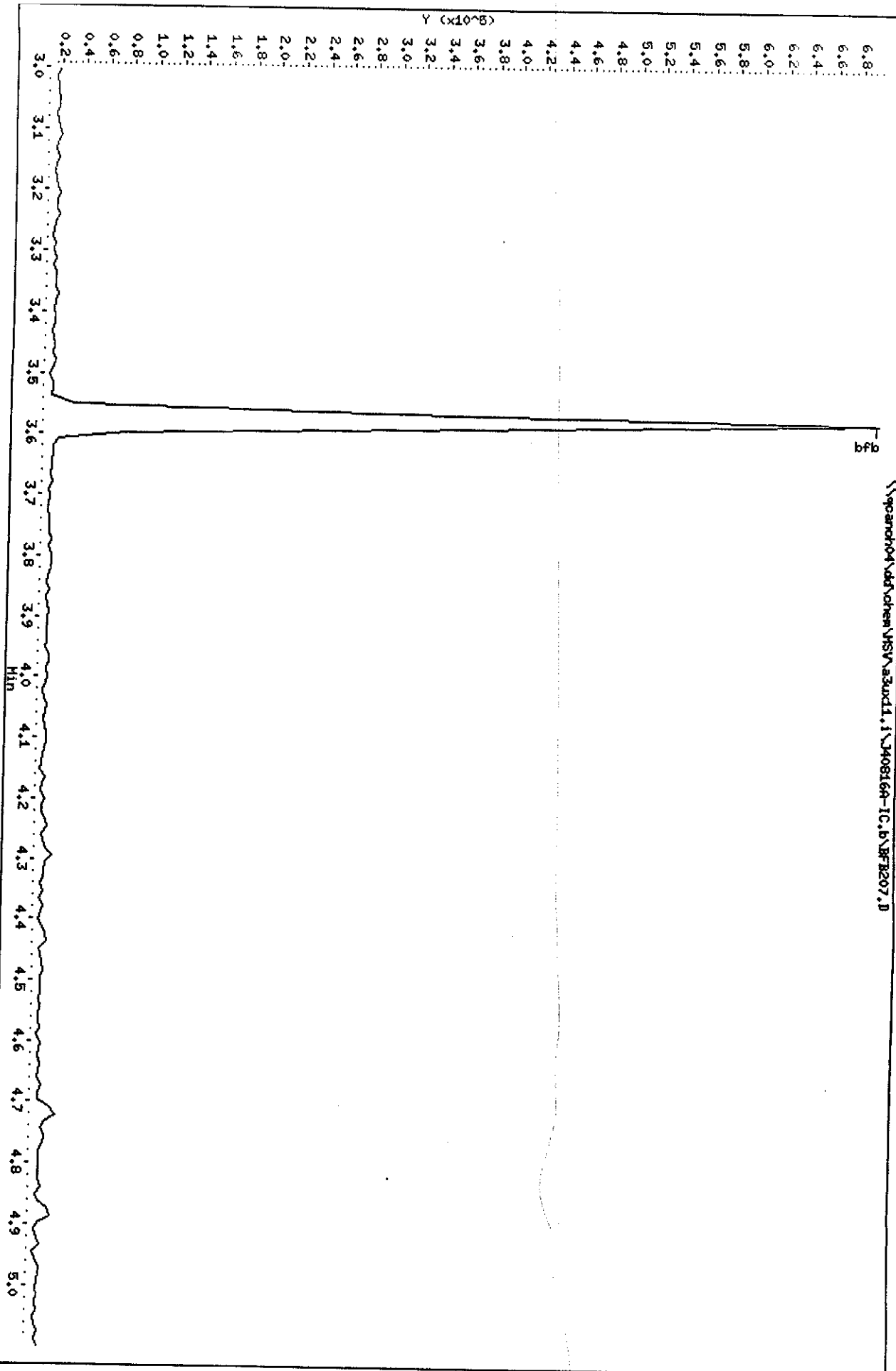
Column Phase: BB624 20H

Instrument: aa3ud1.i

Operator: 43582

Column diameter: 0.18

\\pcaroh04\ddh\chem\NSV\aa3ud1.i\J40816A-IC.b\BFB207.D



Date : 14-SEP-2004 13:21

Client ID: 5ONG BFB

Instrument: z3ux11.i

Sample Info: BFB232

Volume Injected (uL): 1.0

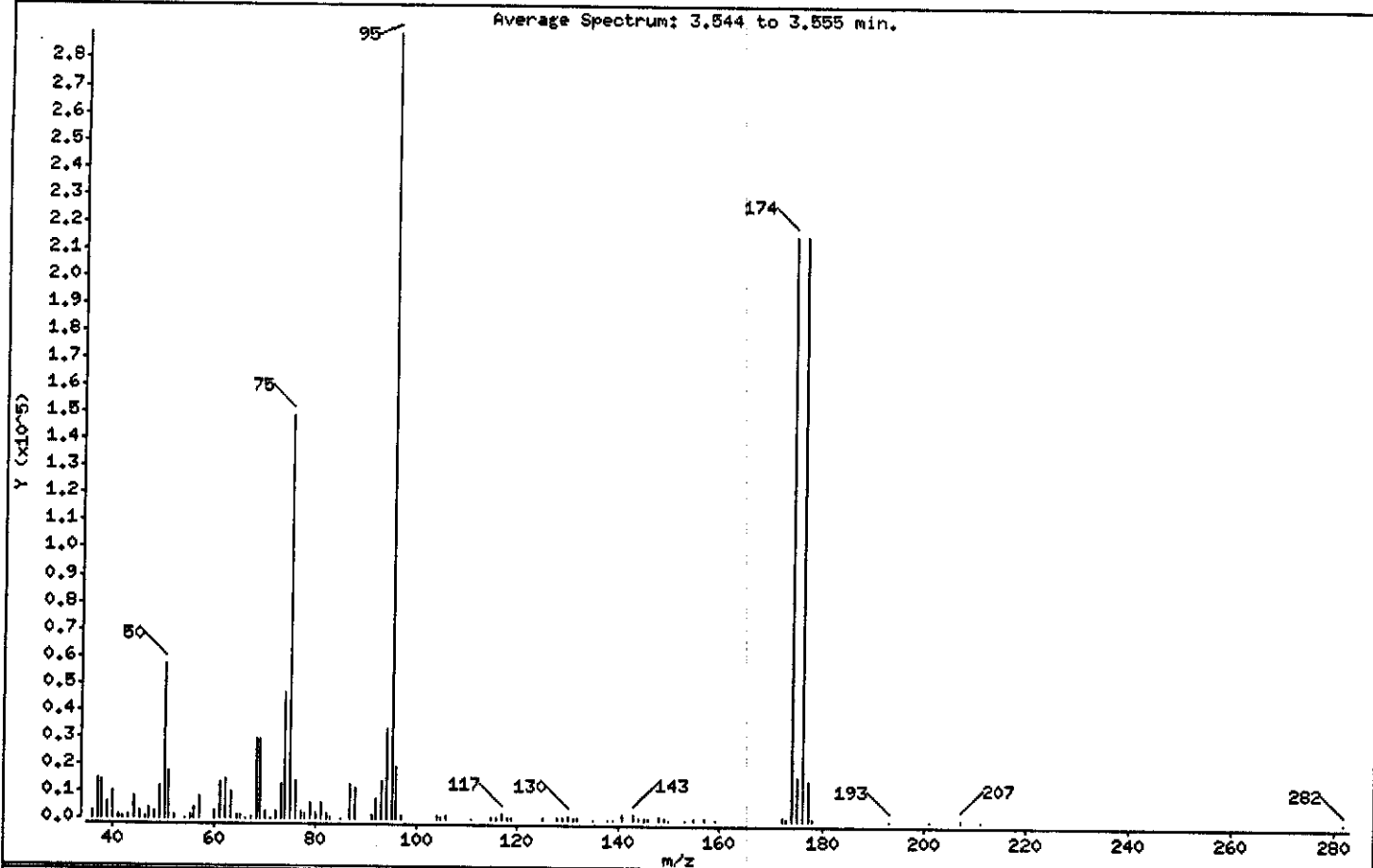
Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

1 bfb

Average Spectrum: 3.544 to 3.555 min.



| m/e | ION ABUNDANCE CRITERIA | X RELATIVE ABUNDANCE |
|-----|------------------------------------|----------------------|
| 95 | Base Peak, 100% relative abundance | 100.00 |
| 50 | 15.00 - 40.00% of mass 95 | 19.83 |
| 75 | 30.00 - 60.00% of mass 95 | 51.55 |
| 96 | 5.00 - 9.00% of mass 95 | 6.64 |
| 173 | Less than 2.00% of mass 174 | 0.21 (< 0.28) |
| 174 | 50.00 - 100.00% of mass 95 | 74.58 |
| 175 | 5.00 - 9.00% of mass 174 | 5.61 (< 7.52) |
| 176 | 95.00 - 101.00% of mass 174 | 74.49 (< 99.87) |
| 177 | 5.00 - 9.00% of mass 176 | 5.17 (< 6.94) |

Date : 14-SEP-2004 13:21

Client ID: 5ONG BFB

Instrument: z3ux11.i

Sample Info: BFB232

Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

Data File: BFB232.D

Spectrum: Average Spectrum: 3.544 to 3.565 min.

Location of Maximum: 95.00

Number of points: 97

| m/z | Y | m/z | Y | m/z | Y | m/z | Y |
|-------|-------|-------|--------|--------|--------|--------|--------|
| 36.00 | 2406 | 64.00 | 1324 | 93.00 | 13655 | 144.00 | 401 |
| 37.00 | 14778 | 65.00 | 1522 | 94.00 | 33528 | 145.00 | 409 |
| 38.00 | 13742 | 66.00 | 263 | 95.00 | 288960 | 146.00 | 877 |
| 39.00 | 5795 | 67.00 | 910 | 96.00 | 19192 | 148.00 | 1075 |
| 40.00 | 10147 | 68.00 | 29456 | 97.00 | 1451 | 149.00 | 479 |
| 41.00 | 1576 | 69.00 | 29344 | 104.00 | 1533 | 150.00 | 313 |
| 42.00 | 737 | 70.00 | 2634 | 105.00 | 833 | 153.00 | 287 |
| 43.00 | 1309 | 71.00 | 324 | 106.00 | 1128 | 155.00 | 668 |
| 44.00 | 8117 | 72.00 | 2356 | 111.00 | 260 | 157.00 | 711 |
| 45.00 | 2441 | 73.00 | 12818 | 115.00 | 375 | 159.00 | 271 |
| 46.00 | 749 | 74.00 | 46784 | 116.00 | 766 | 172.00 | 1628 |
| 47.00 | 3975 | 75.00 | 148928 | 117.00 | 1678 | 173.00 | 603 |
| 48.00 | 2388 | 76.00 | 13979 | 118.00 | 426 | 174.00 | 215488 |
| 49.00 | 12238 | 77.00 | 2377 | 119.00 | 915 | 175.00 | 16212 |
| 50.00 | 57288 | 78.00 | 2058 | 125.00 | 649 | 176.00 | 215232 |
| 51.00 | 17136 | 79.00 | 5934 | 128.00 | 783 | 177.00 | 14940 |
| 52.00 | 1094 | 80.00 | 2304 | 129.00 | 689 | 178.00 | 355 |
| 54.00 | 314 | 81.00 | 6191 | 130.00 | 1263 | 193.00 | 286 |
| 55.00 | 1279 | 82.00 | 1726 | 131.00 | 957 | 201.00 | 259 |
| 56.00 | 4132 | 83.00 | 677 | 132.00 | 360 | 207.00 | 652 |
| 57.00 | 7889 | 85.00 | 327 | 135.00 | 330 | 211.00 | 272 |
| 60.00 | 2651 | 87.00 | 12709 | 138.00 | 255 | 282.00 | 251 |
| 61.00 | 13040 | 88.00 | 11543 | 139.00 | 322 | | |
| 62.00 | 14627 | 91.00 | 1380 | 141.00 | 2064 | | |
| 63.00 | 10230 | 92.00 | 7894 | 143.00 | 2214 | | |

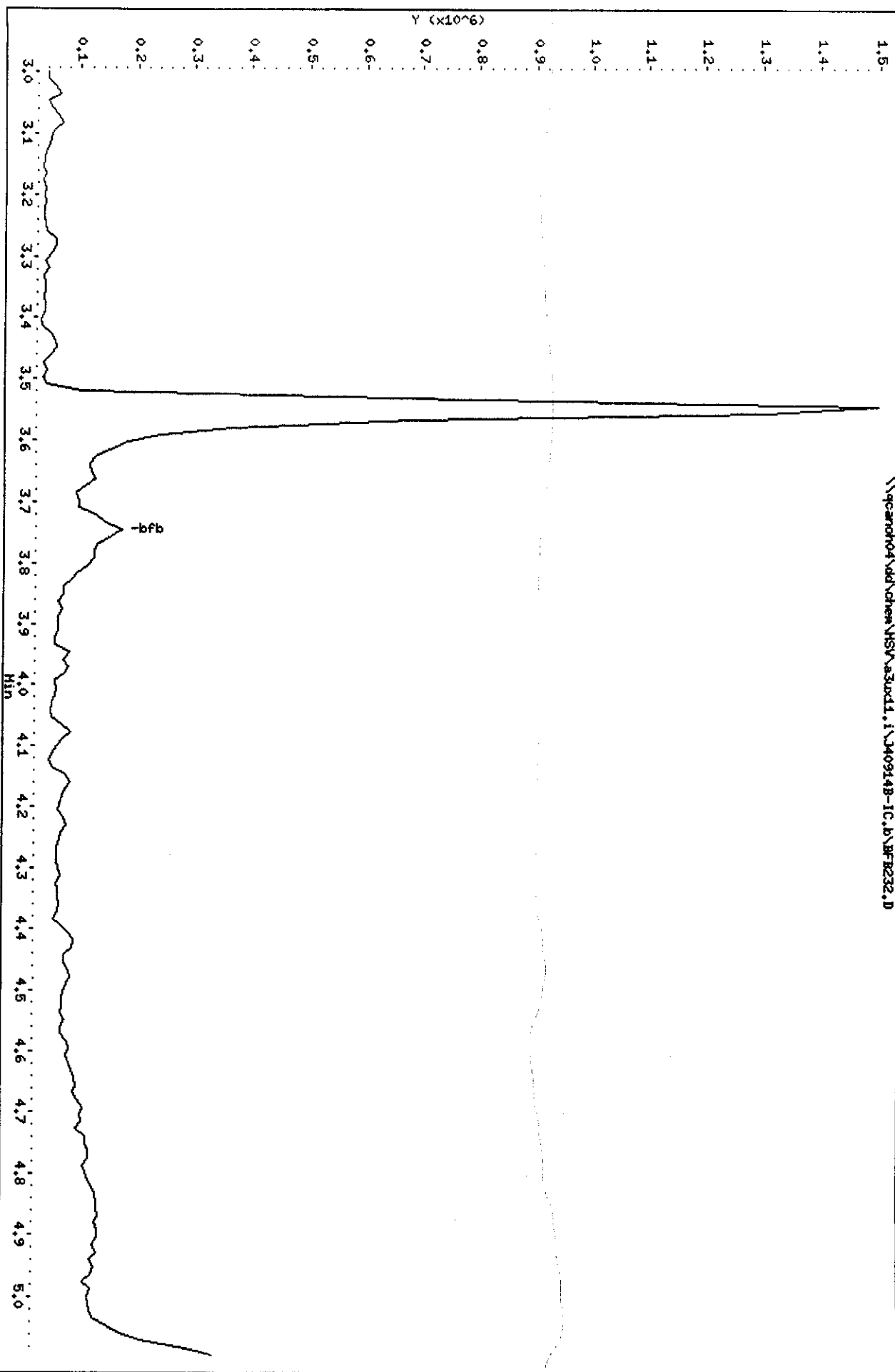
Data File: \\qcarch04\\dd\\chen\\HSV\\a3ux11.i\\J40914B-IC.b\\BF232.D
Date : 14-SEP-2004 13:24
Client ID: SONG EFB

Page 2

Sample Info: BF232
Volume Injected (uL): 1.0
Column phase: DBe24 20H

Instrument: a3ux11.i
Operator: 43562
Column diameter: 0.18

\\qcarch04\\dd\\chen\\HSV\\a3ux11.i\\J40914B-IC.b\\BF232.D



Date : 01-OCT-2004 07:30

Client ID: BONG BFB

Instrument: z3ud11.i

Sample Info:

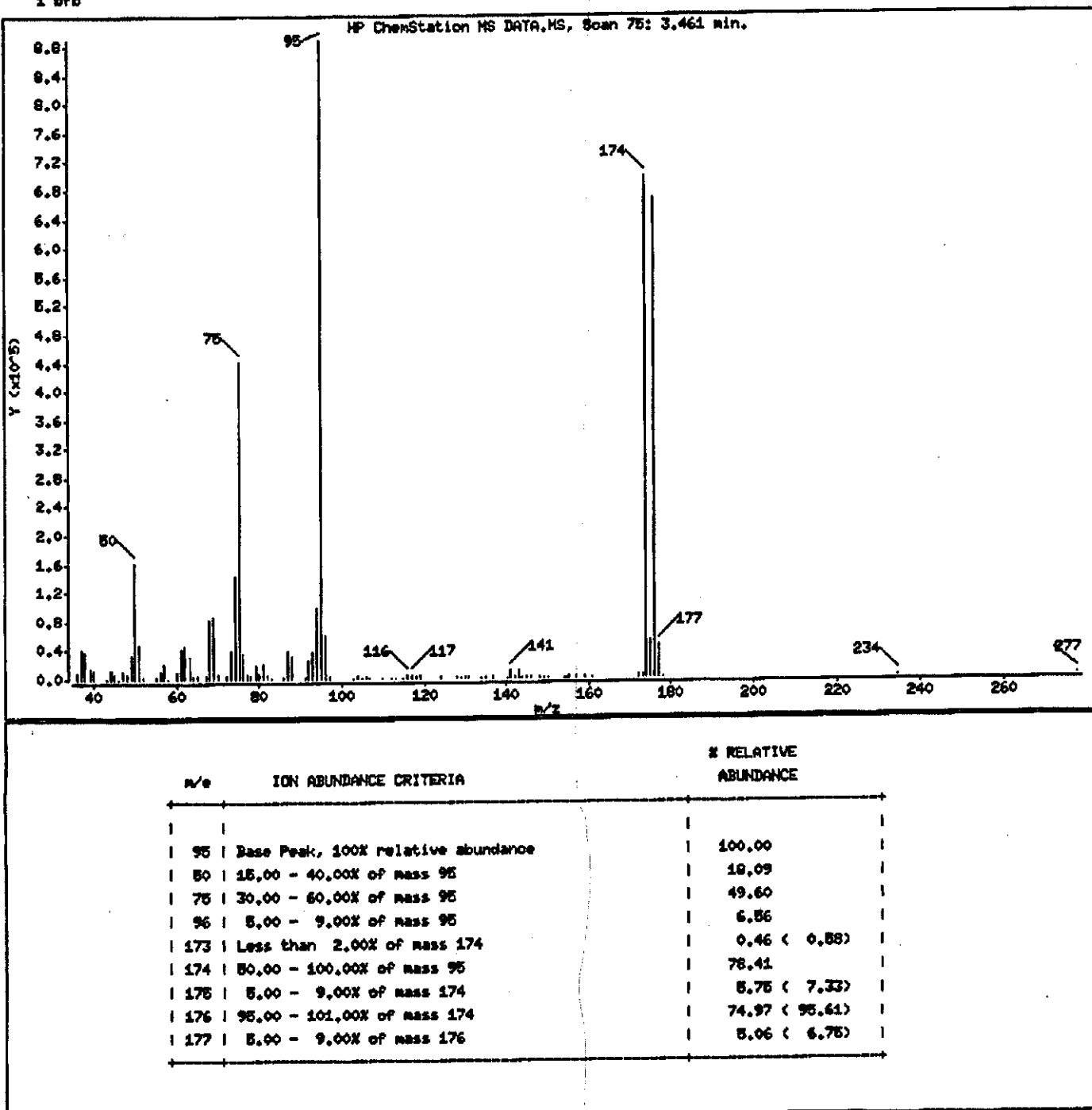
Volume Injected (uL): 1.0

Operator: LE

Column phase: DB624 20M

Column diameter: 0.18

1 bfb



Date : 01-OCT-2004 07:30

Client ID: 50HG BFB

Instrument: z3ud11.i

Sample Info:

Volume Injected (uL): 1.0

Operator: LE

Column phase: ZB624 20H

Column diameter: 0.18

Data File: BFB246.D

Spectrum: HP ChemStation MS DATA.MS, Scan 75; 3.461 min.

Location of Maximum: 95.10

Number of points: 96

| m/z | Y | m/z | Y | m/z | Y | m/z | Y |
|-------|--------|-------|--------|--------|------|--------|--------|
| 36.00 | 7157 | 67.00 | 3180 | 97.10 | 2089 | 143.00 | 9251 |
| 37.10 | 40664 | 68.00 | 82424 | 102.90 | 640 | 143.90 | 601 |
| 38.00 | 38762 | 69.00 | 86368 | 104.00 | 3146 | 145.00 | 1357 |
| 39.10 | 18118 | 70.10 | 7017 | 104.80 | 626 | 146.00 | 1294 |
| 40.00 | 11878 | 72.10 | 4781 | 106.00 | 2870 | 147.90 | 1901 |
| 43.10 | 773 | 73.00 | 38376 | 106.80 | 856 | 149.00 | 794 |
| 44.10 | 11464 | 74.10 | 142592 | 109.80 | 897 | 150.00 | 909 |
| 45.00 | 6628 | 75.10 | 440576 | 111.90 | 747 | 153.90 | 779 |
| 46.10 | 963 | 76.10 | 38288 | 112.90 | 600 | 154.10 | 780 |
| 47.00 | 11093 | 77.00 | 6558 | 114.90 | 945 | 154.40 | 627 |
| 48.00 | 5201 | 77.90 | 3682 | 115.90 | 3334 | 155.00 | 1629 |
| 49.10 | 32868 | 79.00 | 18104 | 117.00 | 4681 | 157.00 | 2027 |
| 50.00 | 160704 | 79.90 | 8743 | 117.90 | 2726 | 159.00 | 1169 |
| 51.00 | 47096 | 80.90 | 20394 | 119.00 | 3898 | 160.90 | 932 |
| 52.00 | 2746 | 82.00 | 4233 | 124.00 | 1115 | 172.00 | 4065 |
| 55.00 | 2516 | 83.00 | 802 | 126.00 | 2912 | 173.00 | 4061 |
| 56.10 | 10704 | 86.00 | 1706 | 129.00 | 1008 | 174.00 | 696576 |
| 57.00 | 20248 | 87.00 | 39520 | 129.90 | 2648 | 176.00 | 81072 |
| 58.10 | 690 | 88.00 | 30904 | 130.90 | 1308 | 176.00 | 665984 |
| 60.00 | 8804 | 91.00 | 2726 | 134.00 | 658 | 177.00 | 44976 |
| 61.00 | 40768 | 92.00 | 24064 | 134.90 | 1135 | 178.00 | 890 |
| 62.00 | 44240 | 93.00 | 36984 | 136.90 | 1553 | 234.60 | 746 |
| 63.10 | 31200 | 94.10 | 97536 | 140.10 | 883 | 277.40 | 866 |
| 64.00 | 3671 | 95.10 | 886320 | 141.00 | 9349 | | |
| 65.00 | 4843 | 96.00 | 88312 | 142.00 | 1166 | | |

Note File: \\pcpanch04\\datachen\\NBS\\n2001.1\\341001A.b\\BF246.b

Date : 01-OCT-2004 07:30

Client ID: 5046 BFB

Sample Info:

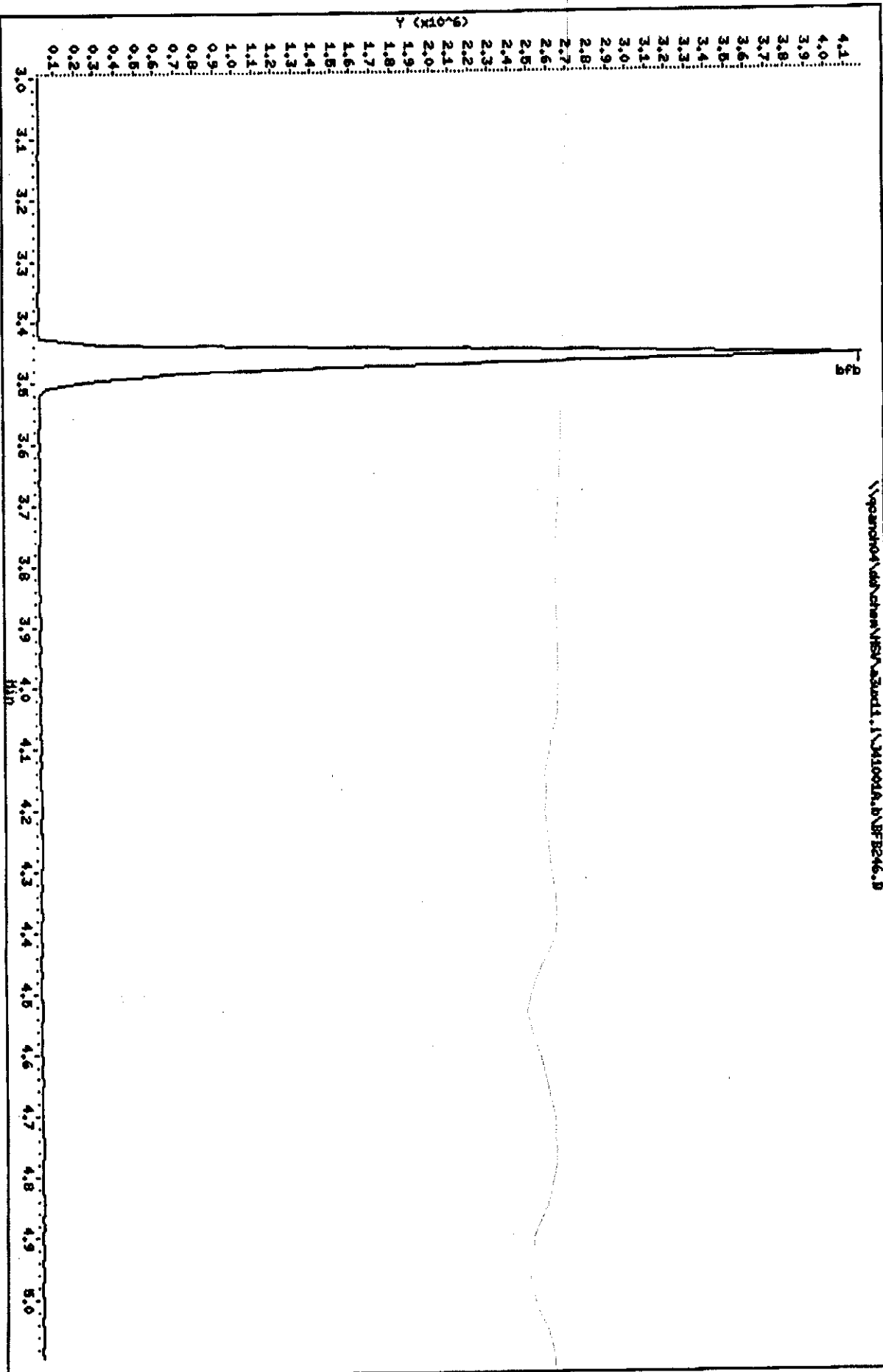
Volume Injected (ul): 1.0

Column phase: DEG24 20H

Instrument: z3001.i

Operator: LE
Column diameter: 0.18

\\pcpanch04\\datachen\\NBS\\n2001.1\\341001A.b\\BF246.b



LABORATORY CONTROL SAMPLE EVALUATION REPORT

GC/MS Volatiles

Client Lot #....: A4I290193 Work Order #....: GRKVP1AC-LCS Matrix.....: WATER
 LCS Lot-Sample#: A4J010000-213 GRKVP1AD-LCSD
 Prep Date.....: 10/01/04 Analysis Date...: 10/01/04
 Prep Batch #....: 4275213
 Dilution Factor: 1 Final Wgt/Vol...: 5 mL
 Initial Wgt/Vol: 5 mL

| <u>PARAMETER</u> | <u>PERCENT RECOVERY</u> | <u>RECOVERY LIMITS</u> | <u>RPD</u> | <u>RPD LIMITS</u> | <u>METHOD</u> |
|---|-------------------------|------------------------|------------|-------------------|---------------|
| Acetone | 71 | (22 - 200) | | | SW846 8260B |
| | 65 | (22 - 200) | 8.3 | (0-95) | SW846 8260B |
| Benzene | 95 | (80 - 116) | | | SW846 8260B |
| | 96 | (80 - 116) | 0.88 | (0-20) | SW846 8260B |
| Bromodichloromethane | 97 | (87 - 130) | | | SW846 8260B |
| | 97 | (87 - 130) | 0.27 | (0-30) | SW846 8260B |
| Bromoform | 106 | (76 - 150) | | | SW846 8260B |
| | 105 | (76 - 150) | 1.1 | (0-30) | SW846 8260B |
| Bromomethane | 59 a | (64 - 129) | | | SW846 8260B |
| | 59 a | (64 - 129) | 0.050 | (0-30) | SW846 8260B |
| 2-Butanone | 80 | (28 - 237) | | | SW846 8260B |
| | 83 | (28 - 237) | 3.7 | (0-65) | SW846 8260B |
| Carbon disulfide | 80 | (73 - 139) | | | SW846 8260B |
| | 78 | (73 - 139) | 1.8 | (0-30) | SW846 8260B |
| Carbon tetrachloride | 84 | (75 - 149) | | | SW846 8260B |
| | 86 | (75 - 149) | 2.3 | (0-30) | SW846 8260B |
| Chlorobenzene | 100 | (76 - 117) | | | SW846 8260B |
| | 102 | (76 - 117) | 2.1 | (0-20) | SW846 8260B |
| Dichlorodifluoromethane | 15 a | (70 - 130) | | | SW846 8260B |
| | 13 a | (70 - 130) | 13 | (0-30) | SW846 8260B |
| 1,1,2-Trichloro- 1,2,2-trifluoroethane | 96 | (70 - 130) | | | SW846 8260B |
| | 95 | (70 - 130) | 1.0 | (0-30) | SW846 8260B |
| Methyl acetate | 94 | (70 - 130) | | | SW846 8260B |
| | 94 | (70 - 130) | 0.37 | (0-30) | SW846 8260B |
| Methyl tert-butyl ether (MTBE) | 81 | (70 - 130) | | | SW846 8260B |
| | 78 | (70 - 130) | 2.8 | (0-30) | SW846 8260B |
| Cyclohexane | 74 | (70 - 130) | | | SW846 8260B |
| | 75 | (70 - 130) | 0.47 | (0-30) | SW846 8260B |
| Methylcyclohexane | 71 | (70 - 130) | | | SW846 8260B |
| | 70 | (70 - 130) | 0.36 | (0-30) | SW846 8260B |
| Dibromochloromethane | 101 | (81 - 138) | | | SW846 8260B |
| | 102 | (81 - 138) | 0.73 | (0-30) | SW846 8260B |
| Isopropylbenzene | 103 | (70 - 130) | | | SW846 8260B |
| | 107 | (70 - 130) | 3.5 | (0-30) | SW846 8260B |

(Continued on next page)

LABORATORY CONTROL SAMPLE EVALUATION REPORT

GC/MS Volatiles

**Client Lot #....: A4I290193 Work Order #....: GRKVP1AC-LCS Matrix.....: WATER
LCS Lot-Sample#: A4J010000-213 GRKVP1AD-LCSD**

| PARAMETER | PERCENT | RECOVERY | RPD | LIMITS | METHOD |
|-----------------------------|-----------------|-----------------|------------|---------------|---------------|
| | RECOVERY | LIMITS | RPD | LIMITS | |
| 1,3-Dichlorobenzene | 93 | (70 - 130) | | | SW846 8260B |
| | 95 | (70 - 130) | 1.4 | (0-30) | SW846 8260B |
| Chloroethane | 75 | (66 - 126) | | | SW846 8260B |
| | 71 | (66 - 126) | 4.5 | (0-30) | SW846 8260B |
| 1,4-Dichlorobenzene | 104 | (70 - 130) | | | SW846 8260B |
| | 102 | (70 - 130) | 1.3 | (0-30) | SW846 8260B |
| 1,2-Dichlorobenzene | 96 | (70 - 130) | | | SW846 8260B |
| | 94 | (70 - 130) | 2.5 | (0-30) | SW846 8260B |
| 1,2,4-Trichloro-benzene | 37 a | (70 - 130) | | | SW846 8260B |
| | 31 a | (70 - 130) | 16 | (0-30) | SW846 8260B |
| Chloroform | 96 | (84 - 128) | | | SW846 8260B |
| | 95 | (84 - 128) | 0.88 | (0-30) | SW846 8260B |
| Chloromethane | 44 a | (48 - 123) | | | SW846 8260B |
| | 44 a | (48 - 123) | 0.40 | (0-30) | SW846 8260B |
| 1,2-Dibromo-3-chloropropane | 76 | (70 - 130) | | | SW846 8260B |
| | 72 | (70 - 130) | 5.5 | (0-30) | SW846 8260B |
| 1,2-Dibromoethane | 101 | (70 - 130) | | | SW846 8260B |
| | 102 | (70 - 130) | 1.1 | (0-30) | SW846 8260B |
| 1,1-Dichloroethane | 93 | (86 - 123) | | | SW846 8260B |
| | 93 | (86 - 123) | 0.060 | (0-30) | SW846 8260B |
| 1,2-Dichloroethane | 99 | (79 - 136) | | | SW846 8260B |
| | 99 | (79 - 136) | 0.19 | (0-30) | SW846 8260B |
| cis-1,2-Dichloroethene | 91 | (85 - 113) | | | SW846 8260B |
| | 92 | (85 - 113) | 1.3 | (0-30) | SW846 8260B |
| trans-1,2-Dichloroethene | 88 | (79 - 120) | | | SW846 8260B |
| | 91 | (79 - 120) | 4.0 | (0-30) | SW846 8260B |
| 1,1-Dichloroethene | 82 | (63 - 130) | | | SW846 8260B |
| | 86 | (63 - 130) | 5.2 | (0-20) | SW846 8260B |
| 1,2-Dichloroethene (total) | 89 | (82 - 116) | | | SW846 8260B |
| | 92 | (82 - 116) | 2.6 | (0-30) | SW846 8260B |
| 1,2-Dichloropropane | 98 | (82 - 115) | | | SW846 8260B |
| | 100 | (82 - 115) | 2.2 | (0-30) | SW846 8260B |
| cis-1,3-Dichloropropene | 87 | (84 - 130) | | | SW846 8260B |
| | 89 | (84 - 130) | 2.8 | (0-30) | SW846 8260B |
| trans-1,3-Dichloropropene | 83 a | (84 - 130) | | | SW846 8260B |
| | 85 | (84 - 130) | 1.8 | (0-30) | SW846 8260B |

(Continued on next page)

LABORATORY CONTROL SAMPLE EVALUATION REPORT

GC/MS Volatiles

Client Lot #....: A4I290193 Work Order #....: GRKVP1AC-LCS Matrix.....: WATER
 LCS Lot-Sample#: A4J010000-213 GRKVP1AD-LCSD

| <u>PARAMETER</u> | <u>PERCENT RECOVERY</u> | <u>RECOVERY LIMITS</u> | <u>RPD</u> | <u>LIMITS</u> | <u>METHOD</u> |
|---------------------------|-------------------------|------------------------|------------|---------------|---------------|
| Ethylbenzene | 97 | (86 - 116) | | | SW846 8260B |
| | 96 | (86 - 116) | 0.14 | (0-30) | SW846 8260B |
| 2-Hexanone | 86 | (35 - 200) | | | SW846 8260B |
| | 82 | (35 - 200) | 4.6 | (0-52) | SW846 8260B |
| Methylene chloride | 100 | (78 - 118) | | | SW846 8260B |
| | 102 | (78 - 118) | 2.0 | (0-30) | SW846 8260B |
| 4-Methyl-2-pentanone | 111 | (78 - 141) | | | SW846 8260B |
| | 111 | (78 - 141) | 0.43 | (0-32) | SW846 8260B |
| Styrene | 108 | (85 - 117) | | | SW846 8260B |
| | 111 | (85 - 117) | 2.3 | (0-30) | SW846 8260B |
| 1,1,2,2-Tetrachloroethane | 116 | (85 - 118) | | | SW846 8260B |
| | 115 | (85 - 118) | 0.81 | (0-30) | SW846 8260B |
| Tetrachloroethene | 91 | (88 - 113) | | | SW846 8260B |
| | 94 | (88 - 113) | 2.8 | (0-30) | SW846 8260B |
| Toluene | 95 | (74 - 119) | | | SW846 8260B |
| | 96 | (74 - 119) | 1.6 | (0-20) | SW846 8260B |
| 1,1,1-Trichloroethane | 77 a | (78 - 140) | | | SW846 8260B |
| | 77 a | (78 - 140) | 0.76 | (0-30) | SW846 8260B |
| 1,1,2-Trichloroethane | 101 | (83 - 122) | | | SW846 8260B |
| | 100 | (83 - 122) | 1.7 | (0-30) | SW846 8260B |
| Trichloroethene | 92 | (75 - 122) | | | SW846 8260B |
| | 91 | (75 - 122) | 1.6 | (0-20) | SW846 8260B |
| Trichlorofluoromethane | 58 a | (70 - 130) | | | SW846 8260B |
| | 60 a | (70 - 130) | 4.6 | (0-30) | SW846 8260B |
| Vinyl chloride | 52 a | (61 - 120) | | | SW846 8260B |
| | 50 a | (61 - 120) | 4.6 | (0-30) | SW846 8260B |
| Xylenes (total) | 104 | (87 - 116) | | | SW846 8260B |
| | 106 | (87 - 116) | 1.3 | (0-30) | SW846 8260B |

| <u>SURROGATE</u> | <u>PERCENT RECOVERY</u> | <u>RECOVERY LIMITS</u> |
|-----------------------|-------------------------|------------------------|
| Dibromofluoromethane | 103 | (73 - 122) |
| | 103 | (73 - 122) |
| 1,2-Dichloroethane-d4 | 107 | (61 - 128) |
| | 104 | (61 - 128) |
| Toluene-d8 | 101 | (76 - 110) |
| | 102 | (76 - 110) |
| 4-Bromofluorobenzene | 115 | (74 - 116) |
| | 116 | (74 - 116) |

NOTE(S) :

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

a Spiked analyte recovery is outside stated control limits.

LABORATORY CONTROL SAMPLE DATA REPORT

GC/MS Volatiles

Client Lot #....: A4I290193 Work Order #....: GRKVP1AC-LCS Matrix.....: WATER
 LCS Lot-Sample#: A4J010000-213 GRKVP1AD-LCSD
 Prep Date.....: 10/01/04 Analysis Date...: 10/01/04
 Prep Batch #....: 4275213
 Dilution Factor: 1 Final Wgt/Vol..: 5 mL
 Initial Wgt/Vol: 5 mL

| <u>PARAMETER</u> | <u>SPIKE AMOUNT</u> | <u>MEASURED AMOUNT</u> | <u>UNITS</u> | <u>PERCENT RECOVERY</u> | <u>RPD</u> | <u>METHOD</u> |
|---|---------------------|------------------------|--------------|-------------------------|------------|---------------|
| Acetone | 10 | 7.1 | ug/L | 71 | | SW846 8260B |
| | 10 | 6.5 | ug/L | 65 | 8.3 | SW846 8260B |
| Benzene | 10 | 9.5 | ug/L | 95 | | SW846 8260B |
| | 10 | 9.6 | ug/L | 96 | 0.88 | SW846 8260B |
| Bromodichloromethane | 10 | 9.7 | ug/L | 97 | | SW846 8260B |
| | 10 | 9.7 | ug/L | 97 | 0.27 | SW846 8260B |
| Bromoform | 10 | 11 | ug/L | 106 | | SW846 8260B |
| | 10 | 11 | ug/L | 105 | 1.1 | SW846 8260B |
| Bromomethane | 10 | 5.9 a | ug/L | 59 | | SW846 8260B |
| | 10 | 5.9 a | ug/L | 59 | 0.050 | SW846 8260B |
| 2-Butanone | 10 | 8.0 | ug/L | 80 | | SW846 8260B |
| | 10 | 8.3 | ug/L | 83 | 3.7 | SW846 8260B |
| Carbon disulfide | 10 | 8.0 | ug/L | 80 | | SW846 8260B |
| | 10 | 7.8 | ug/L | 78 | 1.8 | SW846 8260B |
| Carbon tetrachloride | 10 | 8.4 | ug/L | 84 | | SW846 8260B |
| | 10 | 8.6 | ug/L | 86 | 2.3 | SW846 8260B |
| Chlorobenzene | 10 | 10 | ug/L | 100 | | SW846 8260B |
| | 10 | 10 | ug/L | 102 | 2.1 | SW846 8260B |
| Dichlorodifluoromethane | 10 | 1.5 a | ug/L | 15 | | SW846 8260B |
| | 10 | 1.3 a | ug/L | 13 | 13 | SW846 8260B |
| 1,1,2-Trichloro- 1,2,2-trifluoroethane | 10 | 9.6 | ug/L | 96 | | SW846 8260B |
| | 10 | 9.5 | ug/L | 95 | 1.0 | SW846 8260B |
| Methyl acetate | 10 | 9.4 | ug/L | 94 | | SW846 8260B |
| | 10 | 9.4 | ug/L | 94 | 0.37 | SW846 8260B |
| Methyl tert-butyl ether (MTBE) | 10 | 8.1 | ug/L | 81 | | SW846 8260B |
| | 10 | 7.8 | ug/L | 78 | 2.8 | SW846 8260B |
| Cyclohexane | 10 | 7.4 | ug/L | 74 | | SW846 8260B |
| | 10 | 7.5 | ug/L | 75 | 0.47 | SW846 8260B |
| Methylcyclohexane | 10 | 7.1 | ug/L | 71 | | SW846 8260B |
| | 10 | 7.0 | ug/L | 70 | 0.36 | SW846 8260B |
| Dibromochloromethane | 10 | 10 | ug/L | 101 | | SW846 8260B |
| | 10 | 10 | ug/L | 102 | 0.73 | SW846 8260B |
| Isopropylbenzene | 10 | 10 | ug/L | 103 | | SW846 8260B |
| | 10 | 11 | ug/L | 107 | 3.5 | SW846 8260B |

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LABORATORY CONTROL SAMPLE DATA REPORT

GC/MS Volatiles

Client Lot #...: A4I290193 Work Order #...: GRKVP1AC-LCS Matrix.....: WATER
 LCS Lot-Sample#: A4J010000-213 GRKVP1AD-LCSD

| PARAMETER | SPIKE | MEASURED | UNITS | PERCENT | RPD | METHOD |
|-----------------------------|--------|----------|-------|----------|-------|-------------|
| | AMOUNT | AMOUNT | | RECOVERY | | |
| 1,3-Dichlorobenzene | 10 | 9.3 | ug/L | 93 | | SW846 8260B |
| | 10 | 9.5 | ug/L | 95 | 1.4 | SW846 8260B |
| Chloroethane | 10 | 7.5 | ug/L | 75 | | SW846 8260B |
| | 10 | 7.1 | ug/L | 71 | 4.5 | SW846 8260B |
| 1,4-Dichlorobenzene | 10 | 10 | ug/L | 104 | | SW846 8260B |
| | 10 | 10 | ug/L | 102 | 1.3 | SW846 8260B |
| 1,2-Dichlorobenzene | 10 | 9.6 | ug/L | 96 | | SW846 8260B |
| | 10 | 9.4 | ug/L | 94 | 2.5 | SW846 8260B |
| 1,2,4-Trichloro-benzene | 10 | 3.7 a | ug/L | 37 | | SW846 8260B |
| | 10 | 3.1 a | ug/L | 31 | 16 | SW846 8260B |
| Chloroform | 10 | 9.6 | ug/L | 96 | | SW846 8260B |
| | 10 | 9.5 | ug/L | 95 | 0.88 | SW846 8260B |
| Chloromethane | 10 | 4.4 a | ug/L | 44 | | SW846 8260B |
| | 10 | 4.4 a | ug/L | 44 | 0.40 | SW846 8260B |
| 1,2-Dibromo-3-chloropropane | 10 | 7.6 | ug/L | 76 | | SW846 8260B |
| | 10 | 7.2 | ug/L | 72 | 5.5 | SW846 8260B |
| 1,2-Dibromoethane | 10 | 10 | ug/L | 101 | | SW846 8260B |
| | 10 | 10 | ug/L | 102 | 1.1 | SW846 8260B |
| 1,1-Dichloroethane | 10 | 9.3 | ug/L | 93 | | SW846 8260B |
| | 10 | 9.3 | ug/L | 93 | 0.060 | SW846 8260B |
| 1,2-Dichloroethane | 10 | 9.9 | ug/L | 99 | | SW846 8260B |
| | 10 | 9.9 | ug/L | 99 | 0.19 | SW846 8260B |
| cis-1,2-Dichloroethene | 10 | 9.1 | ug/L | 91 | | SW846 8260B |
| | 10 | 9.2 | ug/L | 92 | 1.3 | SW846 8260B |
| trans-1,2-Dichloroethene | 10 | 8.8 | ug/L | 88 | | SW846 8260B |
| | 10 | 9.1 | ug/L | 91 | 4.0 | SW846 8260B |
| 1,1-Dichloroethene | 10 | 8.2 | ug/L | 82 | | SW846 8260B |
| | 10 | 8.6 | ug/L | 86 | 5.2 | SW846 8260B |
| 1,2-Dichloroethene (total) | 20 | 18 | ug/L | 89 | | SW846 8260B |
| | 20 | 18 | ug/L | 92 | 2.6 | SW846 8260B |
| 1,2-Dichloropropane | 10 | 9.8 | ug/L | 98 | | SW846 8260B |
| | 10 | 10 | ug/L | 100 | 2.2 | SW846 8260B |
| cis-1,3-Dichloropropene | 10 | 8.7 | ug/L | 87 | | SW846 8260B |
| | 10 | 8.9 | ug/L | 89 | 2.8 | SW846 8260B |
| trans-1,3-Dichloropropene | 10 | 8.3 a | ug/L | 83 | | SW846 8260B |
| | 10 | 8.5 | ug/L | 85 | 1.8 | SW846 8260B |

(Continued on next page)

LABORATORY CONTROL SAMPLE DATA REPORT

GC/MS Volatiles

Client Lot #....: A4I290193 Work Order #....: GRKVP1AC-LCS Matrix.....: WATER
LCS Lot-Sample#: A4J010000-213 GRKVP1AD-LCSD

| PARAMETER | SPIKE | MEASURED | | PERCENT | RPD | METHOD |
|---------------------------|---------------|-----------------|--------------|-----------------|------------|---------------|
| | AMOUNT | AMOUNT | UNITS | RECOVERY | | |
| Ethylbenzene | 10 | 9.7 | ug/L | 97 | | SW846 8260B |
| | 10 | 9.6 | ug/L | 96 | 0.14 | SW846 8260B |
| 2-Hexanone | 10 | 8.6 | ug/L | 86 | | SW846 8260B |
| | 10 | 8.2 | ug/L | 82 | 4.6 | SW846 8260B |
| Methylene chloride | 10 | 10 | ug/L | 100 | | SW846 8260B |
| | 10 | 10 | ug/L | 102 | 2.0 | SW846 8260B |
| 4-Methyl-2-pentanone | 10 | 11 | ug/L | 111 | | SW846 8260B |
| | 10 | 11 | ug/L | 111 | 0.43 | SW846 8260B |
| Styrene | 10 | 11 | ug/L | 108 | | SW846 8260B |
| | 10 | 11 | ug/L | 111 | 2.3 | SW846 8260B |
| 1,1,2,2-Tetrachloroethane | 10 | 12 | ug/L | 116 | | SW846 8260B |
| | 10 | 11 | ug/L | 115 | 0.81 | SW846 8260B |
| Tetrachloroethene | 10 | 9.1 | ug/L | 91 | | SW846 8260B |
| | 10 | 9.4 | ug/L | 94 | 2.8 | SW846 8260B |
| Toluene | 10 | 9.5 | ug/L | 95 | | SW846 8260B |
| | 10 | 9.6 | ug/L | 96 | 1.6 | SW846 8260B |
| 1,1,1-Trichloroethane | 10 | 7.7 a | ug/L | 77 | | SW846 8260B |
| | 10 | 7.7 a | ug/L | 77 | 0.76 | SW846 8260B |
| 1,1,2-Trichloroethane | 10 | 10 | ug/L | 101 | | SW846 8260B |
| | 10 | 10 | ug/L | 100 | 1.7 | SW846 8260B |
| Trichloroethene | 10 | 9.2 | ug/L | 92 | | SW846 8260B |
| | 10 | 9.1 | ug/L | 91 | 1.6 | SW846 8260B |
| Trichlorofluoromethane | 10 | 5.8 a | ug/L | 58 | | SW846 8260B |
| | 10 | 6.0 a | ug/L | 60 | 4.6 | SW846 8260B |
| Vinyl chloride | 10 | 5.2 a | ug/L | 52 | | SW846 8260B |
| | 10 | 5.0 a | ug/L | 50 | 4.6 | SW846 8260B |
| Xylenes (total) | 30 | 31 | ug/L | 104 | | SW846 8260B |
| | 30 | 32 | ug/L | 106 | 1.3 | SW846 8260B |

| SURROGATE | PERCENT | RECOVERY |
|-----------------------|-----------------|-----------------|
| | RECOVERY | LIMITS |
| Dibromofluoromethane | 103 | (73 - 122) |
| | 103 | (73 - 122) |
| 1,2-Dichloroethane-d4 | 107 | (61 - 128) |
| | 104 | (61 - 128) |
| Toluene-d8 | 101 | (76 - 110) |
| | 102 | (76 - 110) |
| 4-Bromofluorobenzene | 115 | (74 - 116) |
| | 116 | (74 - 116) |

NOTE(S):

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

a Spiked analyte recovery is outside stated control limits.

Data File: \\pc1\mohs\mohs\hsn\hsn\2001.1\3410019.b\NRI24279.D

Date : 04-OCT-2001 08:17

Client ID:

G R KVPIAC

Sample Info: CHECK

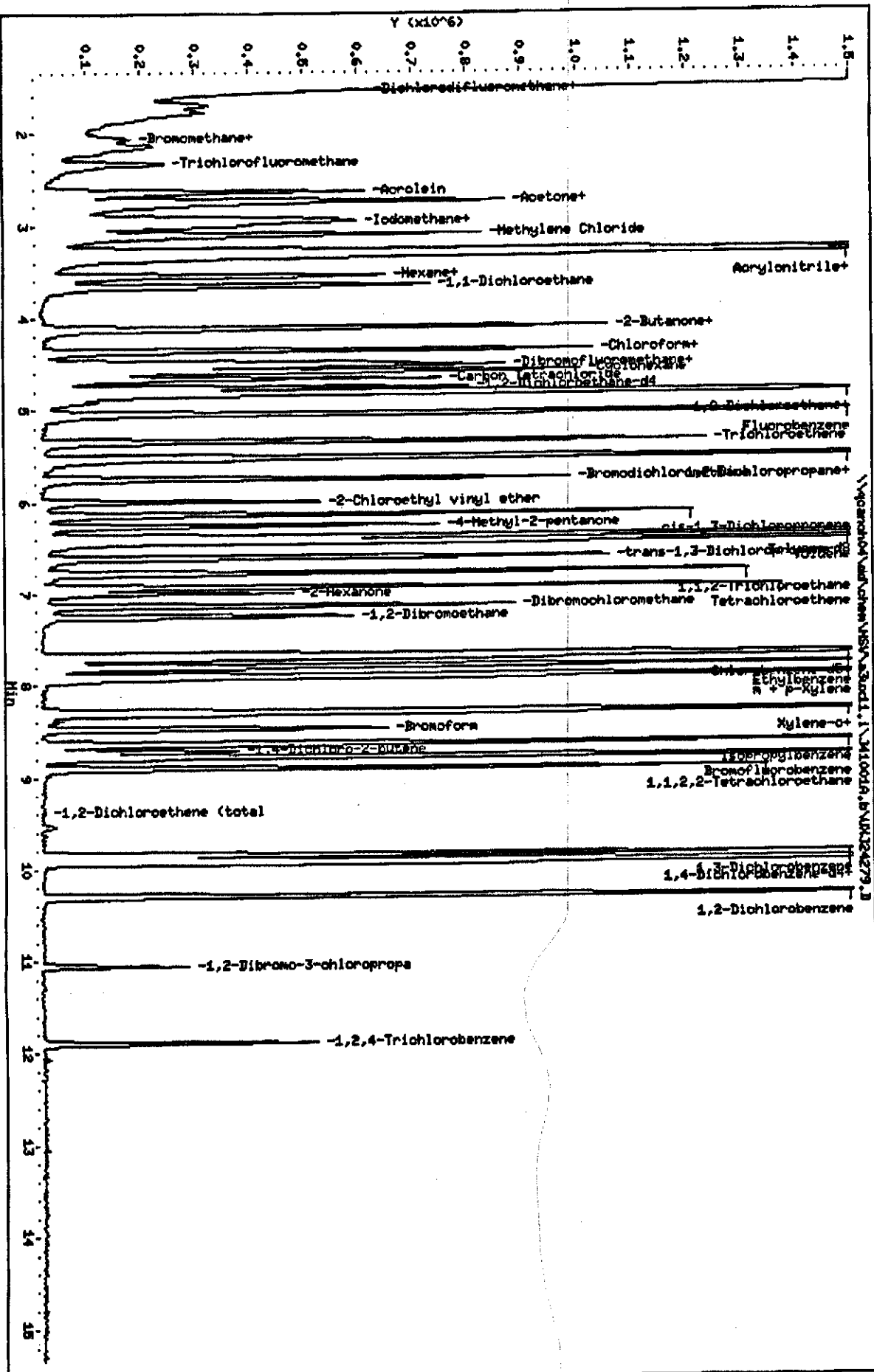
Purge Volume: 5.0

Column Phase: DB-225

Instrument: 4300D.I.

Operator: 4300D.I.

Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24279.D
 Report Date: 04-Oct-2004 09:54

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24279.D
 Lab Smp Id: GRKVP1AC
 Inj Date : 01-OCT-2004 08:47
 Operator : 43582 Inst ID: a3ux11.i
 Smp Info : CHECK
 Misc Info : J41001A,8260LLUX11,2-8260.SUB,43582,3
 Comment :
 Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J41001A.b\8260LLUX11.m
 Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD
 Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
 Als bottle: 3 QC Sample: METHSPIKE
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: 2-8260.SUB
 Target Version: 4.04
 Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|-----------|---------------------------|----------------|-------|---------------|---------|----------|-----------------|---------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) | FINAL (ug/L) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 2028404 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1648603 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 862434 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 487369 | 51.6061 | 10.321 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 691968 | 53.2846 | 10.657 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 2000858 | 50.5191 | 10.104 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 965424 | 57.3693 | 11.474 | |
| \$ | 8 Dichlorodifluoromethane | 85 | 1.639 | 1.639 (0.305) | 80810 | 7.62037 | 1.524 | |
| 9 | Chloromethane | 50 | 1.692 | 1.692 (0.336) | 416864 | 22.0168 | 4.403 | |
| 10 | Vinyl Chloride | 62 | 1.775 | 1.775 (0.352) | 337591 | 26.0658 | 5.213 | |
| 11 | Bromomethane | 94 | 2.059 | 2.059 (0.409) | 182109 | 29.6444 | 5.929 | |
| 12 | Chloroethane | 64 | 2.142 | 2.142 (0.425) | 355258 | 37.3233 | 7.465 | |
| 13 | Trichlorofluoromethane | 101 | 2.320 | 2.320 (0.460) | 390454 | 28.7629 | 5.752 | |
| 15 | Acrolein | 86 | 2.627 | 2.627 (0.521) | 908845 | 717.295 | 143.46 | |
| 16 | Acetone | 43 | 2.745 | 2.745 (0.545) | 176125 | 38.2824 | 7.056 | |
| 17 | 1,1-Dichloroethene | 96 | 2.734 | 2.734 (0.542) | 372031 | 41.0125 | 8.202 | |
| 18 | Freon-113 | 151 | 2.745 | 2.745 (0.545) | 272178 | 47.9321 | 9.586 | |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | |
|---------------------------------|-----------|----------------|-------|---------------|------------------------|-----------|---------|
| | | MASS | RT | EXP RT | REL RT | ON-COLUMN | FINAL |
| | | | | | | (ng) | (ug/L) |
| 19 Iodomethane | | 142 | 2.864 | 2.864 (0.568) | 12358 | 0.92171 | 0.1843 |
| 20 Carbon Disulfide | | 76 | 2.923 | 2.923 (0.580) | 1352862 | 39.9258 | 7.985 |
| 21 Methylene Chloride | | 84 | 3.100 | 3.100 (0.615) | 642493 | 50.1678 | 10.034 |
| 22 Acetonitrile | | 41 | 2.958 | 2.958 (0.587) | 730164 | 606.520 | 121.30 |
| 23 Acrylonitrile | | 53 | 3.278 | 3.278 (0.650) | 2084279 | 556.756 | 111.35 |
| 24 Methyl tert-butyl ether | | 73 | 3.313 | 3.325 (0.657) | 1172004 | 40.3005 | 8.060 |
| 25 trans-1,2-Dichloroethene | | 96 | 3.325 | 3.325 (0.660) | 471679 | 43.8322 | 8.766 |
| 26 Hexane | | 86 | 3.550 | 3.550 (0.704) | 68057 | 39.2717 | 7.854 |
| 27 Vinyl acetate | | 43 | 3.538 | 3.680 (0.702) | 233521 | 13.5614 | 2.712 |
| 28 1,1-Dichloroethane | | 63 | 3.645 | 3.657 (0.723) | 916463 | 46.6858 | 9.337 |
| 29 tert-Butyl Alcohol | | 59 | 3.018 | 3.171 (0.599) | 32050 | 40.6231 | 8.125 |
| 30 2-Butanone | | 43 | 4.094 | 4.106 (0.812) | 222313 | 40.0147 | 8.003 |
| M 31 1,2-Dichloroethane (total) | | 96 | | | 985849 | 89.3079 | 17.862 |
| 32 cis-1,2-dichloroethene | | 96 | 4.106 | 4.106 (0.815) | 514170 | 45.4757 | 9.095 |
| 33 2,2-Dichloropropane | | 77 | | | Compound Not Detected. | | |
| 34 Bromochloromethane | | 128 | | | Compound Not Detected. | | |
| 35 Chloroform | | 83 | 4.355 | 4.355 (0.864) | 950107 | 48.1466 | 9.629 |
| 36 Tetrahydrofuran | | 42 | 4.343 | 4.343 (0.862) | 8083 | 2.78556 | 0.5571 |
| 37 1,1,1-Trichloroethane | | 97 | 4.520 | 4.520 (0.897) | 575963 | 38.6460 | 7.729 |
| 38 1,1-Dichloropropene | | 75 | | | Compound Not Detected. | | |
| 39 Carbon Tetrachloride | | 117 | 4.662 | 4.662 (0.925) | 493717 | 42.1908 | 8.438 |
| 40 1,2-Dichloroethane | | 62 | 4.816 | 4.816 (0.955) | 799696 | 49.5979 | 9.920 |
| 41 Benzene | | 78 | 4.816 | 4.828 (0.955) | 2214770 | 47.4253 | 9.485 |
| 42 Trichloroethene | | 130 | 5.349 | 5.349 (1.061) | 490077 | 45.9813 | 9.196 |
| 43 1,2-Dichloropropane | | 63 | 5.526 | 5.526 (1.096) | 561886 | 48.7745 | 9.755 |
| 44 1,4-Dioxane | | 88 | | | Compound Not Detected. | | |
| 45 Dibromomethane | | 93 | | | Compound Not Detected. | | |
| 46 Bromodichloromethane | | 83 | 5.751 | 5.751 (1.141) | 737636 | 48.6747 | 9.735 |
| 47 2-Chloroethyl vinyl ether | | 63 | 6.000 | 5.999 (1.190) | 269224 | 41.5629 | 8.312 |
| 48 cis-1,3-Dichloropropene | | 75 | 6.130 | 6.130 (1.216) | 806134 | 43.4760 | 8.695 |
| 49 4-Methyl-2-pentanone | | 43 | 6.260 | 6.248 (1.242) | 551835 | 55.3180 | 11.064 |
| 50 Toluene | | 91 | 6.437 | 6.437 (0.838) | 2331449 | 47.3523 | 9.470 |
| 51 trans-1,3-Dichloropropene | | 75 | 6.615 | 6.615 (0.861) | 735971 | 41.6912 | 8.338 |
| 52 Ethyl Methacrylate | | 69 | | | Compound Not Detected. | | |
| 53 1,1,2-Trichloroethane | | 97 | 6.781 | 6.780 (0.883) | 506497 | 50.7152 | 10.143 |
| 54 1,3-Dichloropropane | | 76 | | | Compound Not Detected. | | |
| 55 Tetrachloroethene | | 164 | 6.934 | 6.934 (0.903) | 364236 | 45.6614 | 9.132 |
| 56 2-Hexanone | | 43 | 6.994 | 6.993 (0.911) | 342421 | 42.9508 | 8.590 |
| 57 Dibromochloromethane | | 129 | 7.136 | 7.135 (0.929) | 539785 | 50.6733 | 10.135 |
| 58 1,2-Dibromoethane | | 107 | 7.254 | 7.254 (0.945) | 498253 | 50.6351 | 10.127 |
| 59 Chlorobenzene | | 112 | 7.703 | 7.703 (1.003) | 1596178 | 50.0684 | 10.014 |
| 60 1,1,1,2-Tetrachloroethane | | 131 | | | Compound Not Detected. | | |
| 61 Ethylbenzene | | 106 | 7.798 | 7.798 (1.015) | 777072 | 48.2673 | 9.653 |
| 62 m + p-Xylene | | 106 | 7.905 | 7.905 (1.029) | 2191226 | 105.675 | 21.135 |
| M 63 Xylanes (total) | | 106 | 8.283 | 8.283 (1.079) | 3223685 | 156.394 | 31.279 |
| 64 Xylene-o | | 106 | 8.295 | 8.295 (1.080) | 1967520 | 54.0491 | 10.810 |
| 65 Styrene | | 104 | | | | | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24279.D
 Report Date: 04-Oct-2004 09:54

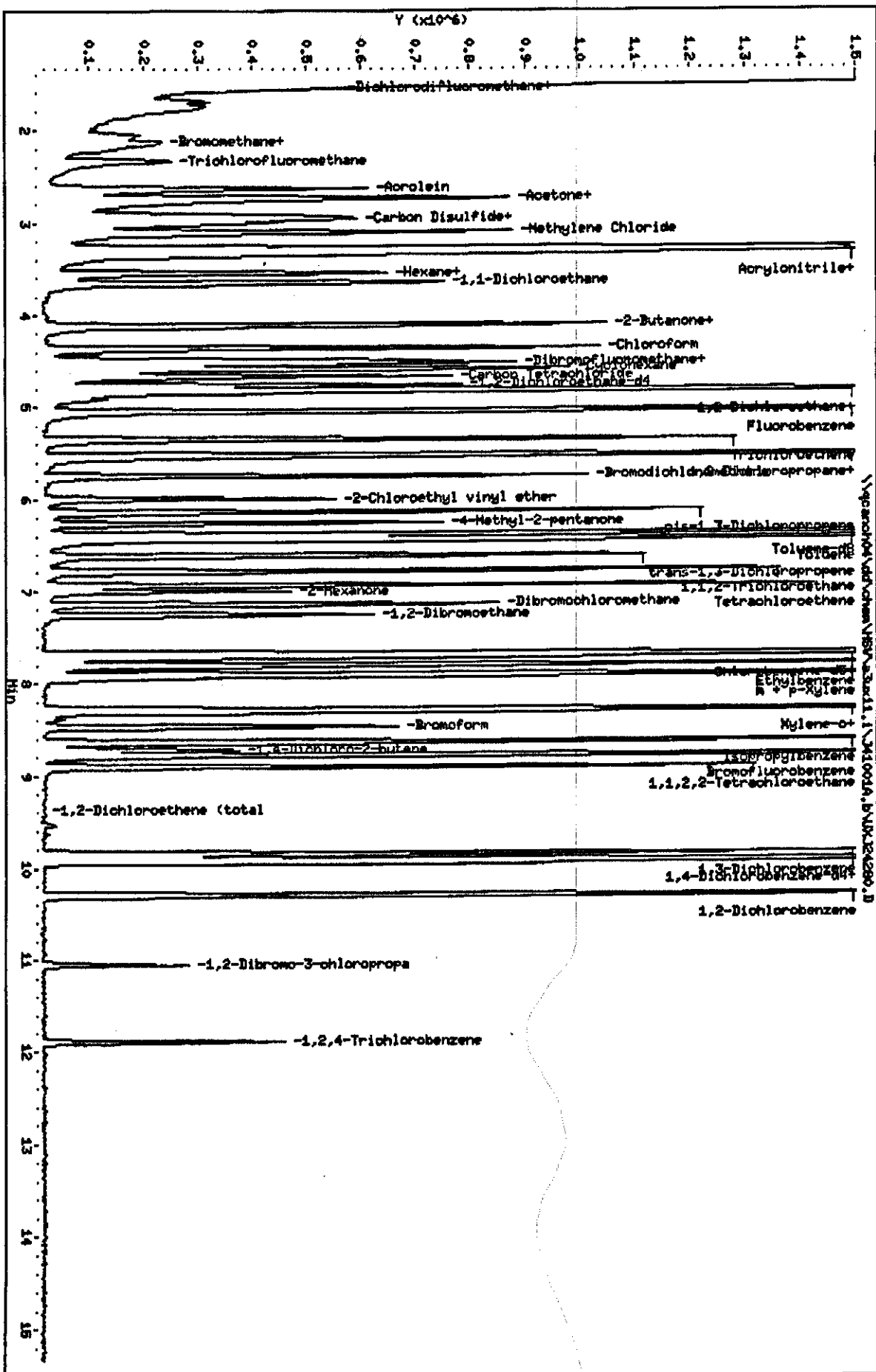
| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | | |
|--------------------------------|-----------|------|----------------|------------------------|--------|----------|----------------|--------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) | FINAL (ug/L) |
| 66 Bromoform | | 173 | 8.473 | 8.473 (1.103) | 1.103 | 384504 | 53.1039 | 10.621 |
| 67 Isopropylbenzene | | 105 | 8.626 | 8.626 (1.123) | 1.123 | 2332276 | 51.4425 | 10.288 |
| 68 1,1,2,2-Tetrachloroethane | | 83 | 8.899 | 8.899 (0.898) | 0.898 | 750776 | 57.7497 | 11.550 |
| 69 1,4-Dichloro-2-butene | | 53 | 8.709 | 8.958 (0.879) | 0.879 | 8679 | 2.02900 | 0.4058 |
| 70 1,2,3-Trichloropropane | | 110 | | Compound Not Detected. | | | | |
| 71 Bromobenzene | | 156 | | Compound Not Detected. | | | | |
| 72 n-Propylbenzene | | 120 | | Compound Not Detected. | | | | |
| 73 2-Chlorotoluene | | 126 | | Compound Not Detected. | | | | |
| 74 1,3,5-Trimethylbenzene | | 105 | | Compound Not Detected. | | | | |
| 75 4-Chlorotoluene | | 126 | | Compound Not Detected. | | | | |
| 76 tert-Butylbenzene | | 119 | | Compound Not Detected. | | | | |
| 77 1,2,4-Trimethylbenzene | | 105 | | Compound Not Detected. | | | | |
| 78 sec-Butylbenzene | | 105 | | Compound Not Detected. | | | | |
| 79 4-Isopropyltoluene | | 119 | | Compound Not Detected. | | | | |
| 80 1,3-Dichlorobenzene | | 146 | 9.845 | 9.845 (0.994) | 1.000 | 1107989 | 46.6417 | 9.328 |
| 81 1,4-Dichlorobenzene | | 146 | 9.928 | 9.928 (1.002) | 1.002 | 1286195 | 51.7512 | 10.350 |
| 82 n-Butylbenzene | | 91 | | Compound Not Detected. | | | | |
| 83 1,2-Dichlorobenzene | | 146 | 10.295 | 10.295 (1.039) | 1.039 | 1125134 | 48.0879 | 9.618 |
| 84 1,2-Dibromo-3-chloropropane | | 157 | 11.052 | 11.052 (1.116) | 1.116 | 83544 | 37.9488 | 7.590 |
| 85 1,2,4-Trichlorobenzene | | 180 | 11.892 | 11.892 (1.201) | 1.201 | 182230 | 18.2942 | 3.659 |
| 86 Hexachlorobutadiene | | 225 | | Compound Not Detected. | | | | |
| 87 Naphthalene | | 126 | | Compound Not Detected. | | | | |
| 88 1,2,3-Trichlorobenzene | | 180 | | Compound Not Detected. | | | | |
| 90 Cyclohexane | | 56 | 4.580 | 4.580 (0.908) | 0.908 | 554896 | 37.1851 | 7.437 |
| 143 Methyl Acetate | | 43 | 3.006 | 3.006 (0.596) | 0.596 | 392704 | 47.0644 | 9.413 |
| 144 Methylcyclohexane | | 83 | 5.526 | 5.526 (1.096) | 1.096 | 442476 | 35.3348 | 7.067 |
| 141 1,3,5-Trichlorobenzene | | 180 | | Compound Not Detected. | | | | |

Date File: \\pcanon4\chem\HSV\30x11.1\J41001A.b\UK24280.D
Date : 01-OCT-2004 09:09
Client ID: G R K V P A D

Sample Info: DECK
Purge Volume: 8.0
Column Phase: DB624

Instrument: Z30x11.1

Operator: 43562
Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24280.D
Report Date: 04-Oct-2004 09:55

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24280.D
Lab Smp Id: grkvplad
Inj Date : 01-OCT-2004 09:09
Operator : 43582 Inst ID: a3ux11.i
Smp Info : CHECK
Misc Info : J41001A,8260LLUX11,2-8260.SUB,43582,3
Comment :
Method : \\QCANOH04\dd\chem\MSV\ a3ux11.i\J41001A.b\8260LLUX11.m
Meth Date : 04-Oct-2004 09:54 evans1 Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 4 QC Sample: METHSPIKE
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|-----------|---------------------------|----------------|-------|------------------------|---------|----------|--------|---------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) | (ug/L) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 2044371 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1659474 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 878061 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 492110 | 51.7011 | 10.340 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 682384 | 52.1362 | 10.427 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 2040953 | 51.1938 | 10.239 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 978570 | 57.7695 | 11.554 | |
| \$ | 8 Dichlorodifluoromethane | 85 | 1.550 | 1.550 (0.308) | 71848 | 6.72234 | 1.344 | |
| 9 | Chloromethane | 50 | 1.692 | 1.704 (0.336) | 421845 | 22.1059 | 4.421 | |
| 10 | Vinyl Chloride | 62 | 1.787 | 1.787 (0.355) | 324827 | 24.8844 | 4.977 | |
| 11 | Bromomethane | 94 | 2.047 | 2.059 (0.406) | 183446 | 29.6288 | 5.926 | |
| 12 | Chloroethane | 64 | 2.130 | 2.142 (0.423) | 342307 | 35.6618 | 7.136 | |
| 13 | Trichlorofluoromethane | 101 | 2.331 | 2.331 (0.462) | 411887 | 30.1048 | 6.021 | |
| 15 | Acrolein | 56 | 2.639 | 2.639 (0.524) | 912057 | 714.208 | 142.84 | |
| 16 | Acetone | 43 | 2.745 | 2.745 (0.545) | 165247 | 32.4613 | 6.492 | |
| 17 | 1,1-Dichloroethene | 96 | 2.733 | 2.734 (0.542) | 394949 | 43.1989 | 8.640 | |
| 18 | Freon-113 | 151 | 2.757 | 2.745 (0.547) | 271383 | 47.4261 | 9.485 | |
| 19 | Iodomethane | 142 | | Compound Not Detected. | | | | |

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J41001A.b\UXJ24280.D
 Report Date: 04-Oct-2004 09:55

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | |
|---------------------------------|-----------|----------------|-------|---------|------------------------|----------|----------------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) |
| 20 Carbon Disulfide | 76 | 2.923 | 2.923 | (0.580) | 1339829 | 39.2324 | 7.846 |
| 21 Methylene Chloride | 84 | 3.100 | 3.100 | (0.615) | 658693 | 51.1950 | 10.239 |
| 22 Acetonitrile | 41 | 2.958 | 2.958 | (0.587) | 806698 | 664.861 | 132.97 |
| 23 Acrylonitrile | 53 | 3.278 | 3.278 | (0.650) | 2112848 | 559.979 | 112.00 |
| 24 Methyl tert-butyl ether | 73 | 3.325 | 3.325 | (0.660) | 1148360 | 39.1791 | 7.836 |
| 25 trans-1,2-Dichloroethene | 96 | 3.325 | 3.325 | (0.660) | 494599 | 45.6032 | 9.121 |
| 26 Hexane | 86 | 3.550 | 3.550 | (0.704) | 70817 | 40.5232 | 8.105 |
| 27 Vinyl acetate | 43 | 3.538 | 3.680 | (0.702) | 238156 | 13.7226 | 2.744 |
| 28 1,1-Dichloroethane | 63 | 3.645 | 3.657 | (0.723) | 924298 | 46.7172 | 9.343 |
| 29 tert-Butyl Alcohol | 59 | 3.325 | 3.171 | (0.660) | 25962 | 32.6496 | 6.530 |
| 30 2-Butanone | 43 | 4.106 | 4.106 | (0.815) | 232466 | 41.5154 | 8.303 |
| M 31 1,2-Dichloroethene (total) | 96 | | | | 1019505 | 91.6658 | 18.333 |
| 32 cis-1,2-dichloroethene | 96 | 4.106 | 4.106 | (0.815) | 524906 | 46.0627 | 9.212 |
| 33 2,2-Dichloropropane | 77 | | | | Compound Not Detected. | | |
| 34 Bromochloromethane | 128 | | | | Compound Not Detected. | | |
| 35 Chloroform | 83 | 4.355 | 4.355 | (0.864) | 949122 | 47.7211 | 9.544 |
| 36 Tetrahydrofuran | 42 | 4.106 | 4.343 | (0.815) | 13990 | 4.62776 | 0.9256 |
| 37 1,1,1-Trichloroethane | 97 | 4.520 | 4.520 | (0.897) | 576059 | 38.3506 | 7.670 |
| 38 1,1-Dichloropropene | 75 | | | | Compound Not Detected. | | |
| 39 Carbon Tetrachloride | 117 | 4.662 | 4.662 | (0.925) | 509148 | 43.1696 | 8.634 |
| 40 1,2-Dichloroethane | 62 | 4.828 | 4.816 | (0.958) | 807545 | 49.6935 | 9.939 |
| 41 Benzene | 78 | 4.828 | 4.828 | (0.958) | 2251900 | 47.8437 | 9.569 |
| 42 Trichloroethene | 130 | 5.349 | 5.349 | (1.061) | 486173 | 45.2587 | 9.052 |
| 43 1,2-Dichloropropane | 63 | 5.526 | 5.526 | (1.096) | 578821 | 49.8521 | 9.970 |
| 44 1,4-Dioxane | 88 | | | | Compound Not Detected. | | |
| 45 Dibromomethane | 93 | | | | Compound Not Detected. | | |
| 46 Bromodichloromethane | 83 | 5.751 | 5.751 | (1.141) | 741406 | 48.5414 | 9.708 |
| 47 2-Chloroethyl vinyl ether | 63 | 5.999 | 5.999 | (1.190) | 269926 | 41.3459 | 8.269 |
| 48 cis-1,3-Dichloropropene | 75 | 6.129 | 6.130 | (1.216) | 835199 | 44.6918 | 8.938 |
| 49 4-Methyl-2-pentanone | 43 | 6.260 | 6.248 | (1.242) | 858308 | 55.8600 | 11.112 |
| 50 Toluene | 91 | 6.437 | 6.437 | (0.838) | 2384206 | 48.1066 | 9.621 |
| 51 trans-1,3-Dichloropropene | 75 | 6.615 | 6.615 | (0.861) | 754081 | 42.4372 | 8.487 |
| 52 Ethyl Methacrylate | 69 | | | | Compound Not Detected. | | |
| 53 1,1,2-Trichloroethane | 97 | 6.780 | 6.780 | (0.883) | 501075 | 49.8436 | 9.969 |
| 54 1,3-Dichloropropane | 76 | | | | Compound Not Detected. | | |
| 55 Tetrachloroethene | 164 | 6.934 | 6.934 | (0.903) | 377104 | 46.9649 | 9.393 |
| 56 2-Hexanone | 43 | 6.993 | 6.993 | (0.911) | 329258 | 41.0292 | 8.206 |
| 57 Dibromochloromethane | 129 | 7.135 | 7.135 | (0.929) | 847365 | 51.0482 | 10.210 |
| 58 1,2-Dibromoethane | 107 | 7.254 | 7.254 | (0.945) | 507352 | 51.2220 | 10.244 |
| 59 Chlorobenzene | 112 | 7.703 | 7.703 | (1.003) | 1641068 | 51.1393 | 10.228 |
| 60 1,1,1,2-Tetrachloroethane | 131 | | | | Compound Not Detected. | | |
| 61 Ethylbenzene | 106 | 7.798 | 7.798 | (1.016) | 781022 | 48.1948 | 9.639 |
| 62 m + p-Xylene | 106 | 7.904 | 7.905 | (1.029) | 2216784 | 106.207 | 21.241 |
| M 63 Xylenes (total) | 106 | | | | 3287082 | 158.441 | 31.688 |
| 64 Xylene-o | 106 | 8.283 | 8.283 | (1.079) | 1070298 | 52.2337 | 10.447 |
| 65 Styrene | 104 | 8.295 | 8.295 | (1.080) | 2025967 | 55.2901 | 11.058 |
| 66 Bromoform | 173 | 8.472 | 8.473 | (1.103) | 382920 | 52.5387 | 10.508 |

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|--------------------------------|-----------|------|----------------|------------------------|---------|----------|-------------------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 67 Isopropylbenzene | | 105 | 8.638 | 8.626 (1.125) | 2431539 | 53.2806 | 10.656 |
| 68 1,1,2,2-Tetrachloroethane | | 83 | 8.898 | 8.899 (0.898) | 758153 | 57.2793 | 11.456 |
| 69 1,4-Dichloro-2-butene | | 53 | 8.709 | 8.958 (0.879) | 6323 | 1.45190 | 0.2904 |
| 70 1,2,3-Trichloropropane | | 110 | | Compound Not Detected. | | | |
| 71 Bromobenzene | | 156 | | Compound Not Detected. | | | |
| 72 n-Propylbenzene | | 120 | | Compound Not Detected. | | | |
| 73 2-Chlorotoluene | | 126 | | Compound Not Detected. | | | |
| 74 1,3,5-Trimethylbenzene | | 105 | | Compound Not Detected. | | | |
| 75 4-Chlorotoluene | | 126 | | Compound Not Detected. | | | |
| 76 tert-Butylbenzene | | 119 | | Compound Not Detected. | | | |
| 77 1,2,4-Trimethylbenzene | | 105 | | Compound Not Detected. | | | |
| 78 sec-Butylbenzenes | | 105 | | Compound Not Detected. | | | |
| 79 4-Isopropyltoluene | | 119 | | Compound Not Detected. | | | |
| 80 1,3-Dichlorobenzene | | 146 | 9.845 | 9.845 (0.994) | 1143366 | 47.2744 | 9.455 |
| 81 1,4-Dichlorobenzene | | 146 | 9.928 | 9.928 (1.002) | 1292846 | 51.0930 | 10.219 |
| 82 n-Butylbenzene | | 91 | | Compound Not Detected. | | | |
| 83 1,2-Dichlorobenzene | | 146 | 10.295 | 10.295 (1.039) | 1117057 | 46.8930 | 9.379 |
| 84 1,2-Dibromo-3-chloropropane | | 157 | 11.052 | 11.052 (1.116) | 80513 | 35.9211 | 7.184 |
| 85 1,2,4-Trichlorobenzene | | 180 | 11.892 | 11.892 (1.201) | 157424 | 15.5226 | 3.104 |
| 86 Hexachlorobutadiene | | 225 | | Compound Not Detected. | | | |
| 87 Naphthalene | | 128 | | Compound Not Detected. | | | |
| 88 1,2,3-Trichlorobenzene | | 180 | | Compound Not Detected. | | | |
| 98 Cyclohexane | | 56 | 4.579 | 4.580 (0.908) | 561869 | 37.3583 | 7.472 |
| 143 Methyl Acetate | | 43 | 3.006 | 3.006 (0.596) | 397276 | 47.2405 | 9.448 |
| 144 Methylcyclohexane | | 83 | 5.526 | 5.526 (1.096) | 444320 | 35.2049 | 7.041 |
| 141 1,3,5-Trichlorobenzene | | 180 | | Compound Not Detected. | | | |

METHOD BLANK REPORT

GC/MS Volatiles

Client Lot #....: A4I290193
 MB Lot-Sample #: A4J010000-213
 Analysis Date...: 10/01/04
 Dilution Factor: 1

Work Order #....: GRKVP1AA
 Prep Date.....: 10/01/04
 Prep Batch #:....: 4275213
 Initial Wgt/Vol: 5 mL

Matrix.....: WATER

Final Wgt/Vol.: 5 mL

| PARAMETER | RESULT | REPORTING | | |
|------------------------------------|--------|-----------|-------|-------------|
| | | LIMIT | UNITS | METHOD |
| Acetone | ND | 10 | ug/L | SW846 8260B |
| Acetonitrile | ND | 20 | ug/L | SW846 8260B |
| Acrolein | ND | 20 | ug/L | SW846 8260B |
| Acrylonitrile | ND | 20 | ug/L | SW846 8260B |
| Benzene | ND | 1.0 | ug/L | SW846 8260B |
| Bromodichloromethane | ND | 1.0 | ug/L | SW846 8260B |
| Bromoform | ND | 1.0 | ug/L | SW846 8260B |
| Bromomethane | ND | 1.0 | ug/L | SW846 8260B |
| 2-Butanone | ND | 10 | ug/L | SW846 8260B |
| Carbon disulfide | ND | 1.0 | ug/L | SW846 8260B |
| Carbon tetrachloride | ND | 1.0 | ug/L | SW846 8260B |
| Chlorobenzene | ND | 1.0 | ug/L | SW846 8260B |
| Chloroprene | ND | 2.0 | ug/L | SW846 8260B |
| Dibromochloromethane | ND | 1.0 | ug/L | SW846 8260B |
| Chloroethane | ND | 1.0 | ug/L | SW846 8260B |
| Chloroform | ND | 1.0 | ug/L | SW846 8260B |
| Chloromethane | 0.21 J | 1.0 | ug/L | SW846 8260B |
| 3-Chloropropene | ND | 2.0 | ug/L | SW846 8260B |
| 1,2-Dibromo-3-chloropropane (DBCP) | ND | 2.0 | ug/L | SW846 8260B |
| 1,2-Dibromoethane | ND | 1.0 | ug/L | SW846 8260B |
| Dibromomethane | ND | 1.0 | ug/L | SW846 8260B |
| trans-1,4-Dichloro-2-butene | ND | 1.0 | ug/L | SW846 8260B |
| 1,1-Dichloroethane | ND | 1.0 | ug/L | SW846 8260B |
| 1,2-Dichloroethane | ND | 1.0 | ug/L | SW846 8260B |
| cis-1,2-Dichloroethene | ND | 1.0 | ug/L | SW846 8260B |
| trans-1,2-Dichloroethene | ND | 1.0 | ug/L | SW846 8260B |
| 1,1-Dichloroethene | ND | 1.0 | ug/L | SW846 8260B |
| 1,2-Dichloroethene (total) | ND | 2.0 | ug/L | SW846 8260B |
| Dichlorofluoromethane | ND | 2.0 | ug/L | SW846 8260B |
| 1,2-Dichloropropane | ND | 1.0 | ug/L | SW846 8260B |
| cis-1,3-Dichloropropene | ND | 1.0 | ug/L | SW846 8260B |
| trans-1,3-Dichloropropene | ND | 1.0 | ug/L | SW846 8260B |
| 1,4-Dioxane | ND | 50 | ug/L | SW846 8260B |
| Ethylbenzene | ND | 1.0 | ug/L | SW846 8260B |
| Ethyl methacrylate | ND | 1.0 | ug/L | SW846 8260B |
| 2-Hexanone | ND | 10 | ug/L | SW846 8260B |
| Iodomethane | ND | 1.0 | ug/L | SW846 8260B |
| Isobutanol | ND | 50 | ug/L | SW846 8260B |

(Continued on next page)

METHOD BLANK REPORT

GC/MS Volatiles

Client Lot #....: A4I290193

Work Order #....: GRKVP1AA

Matrix.....: WATER

| <u>PARAMETER</u> | <u>RESULT</u> | <u>REPORTING</u> | | |
|---------------------------|---------------|------------------|--------------|---------------|
| | | <u>LIMIT</u> | <u>UNITS</u> | <u>METHOD</u> |
| Methacrylonitrile | ND | 2.0 | ug/L | SW846 8260B |
| Methylene chloride | ND | 1.0 | ug/L | SW846 8260B |
| Methyl methacrylate | ND | 2.0 | ug/L | SW846 8260B |
| 4-Methyl-2-pentanone | ND | 10 | ug/L | SW846 8260B |
| Propionitrile | ND | 4.0 | ug/L | SW846 8260B |
| Styrene | ND | 1.0 | ug/L | SW846 8260B |
| 1,1,1,2-Tetrachloroethane | ND | 1.0 | ug/L | SW846 8260B |
| 1,1,2,2-Tetrachloroethane | ND | 1.0 | ug/L | SW846 8260B |
| Tetrachloroethene | ND | 1.0 | ug/L | SW846 8260B |
| Toluene | ND | 1.0 | ug/L | SW846 8260B |
| 1,1,1-Trichloroethane | ND | 1.0 | ug/L | SW846 8260B |
| 1,1,2-Trichloroethane | ND | 1.0 | ug/L | SW846 8260B |
| Trichloroethene | ND | 1.0 | ug/L | SW846 8260B |
| Trichlorofluoromethane | ND | 1.0 | ug/L | SW846 8260B |
| 1,2,3-Trichloropropane | ND | 1.0 | ug/L | SW846 8260B |
| Vinyl acetate | ND | 2.0 | ug/L | SW846 8260B |
| Vinyl chloride | ND | 1.0 | ug/L | SW846 8260B |
| Xylenes (total) | ND | 2.0 | ug/L | SW846 8260B |

| <u>SURROGATE</u> | <u>PERCENT</u> | <u>RECOVERY</u> |
|-----------------------|-----------------|-----------------|
| | <u>RECOVERY</u> | <u>LIMITS</u> |
| Dibromofluoromethane | 115 | (73 - 122) |
| 1,2-Dichloroethane-d4 | 114 | (61 - 128) |
| Toluene-d8 | 88 | (76 - 110) |
| 4-Bromofluorobenzene | 78 | (74 - 116) |

NOTE(S) :

Calculations are performed before rounding to avoid round-off errors in calculated results.

J Estimated result. Result is less than RL.

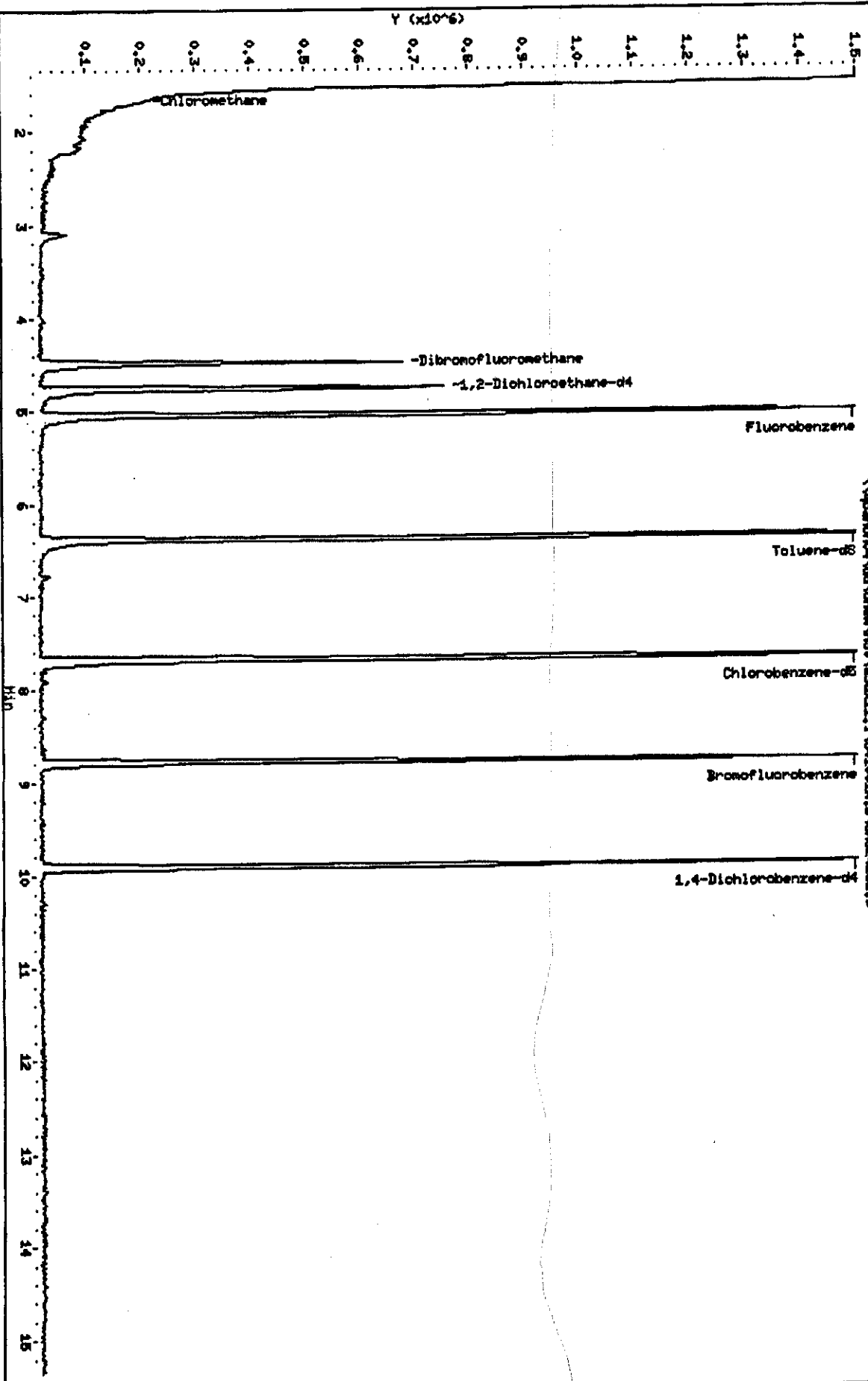
Data File: \\pcando04\detach\HSVA3nd1.1\N41001A.J\N41001A.D
Date : 01-DEC-2004 09:13
Client ID: GRRUPIAM

Sample Info: VALK
Purge Volume: 5.0
Column phase: RK24

Instrument: a3nd1.i

Operator: 43582
Column diameter: 0.18

\\pcando04\detach\HSVA3nd1.1\N41001A.J\N41001A.D



Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24281.D
Report Date: 04-Oct-2004 09:56

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\ a3ux11.i\J41001A.b\UXJ24281.D

Lab Smp Id: grkvplaa

Inj Date : 01-OCT-2004 09:32

Inst ID: a3ux11.i

Operator : 43582

Smp Info : VBLK

Misc Info : J41001A,8260LLUX11,,43582,3,,BLANK,,0

Comment :

Method : \\QCANOH04\dd\chem\MSV\ a3ux11.i\J41001A.b\8260LLUX11.m

Meth Date : 04-Oct-2004 09:54 evansl Quant Type: ISTD

Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D

Als bottle: 5 QC Sample: BLANK

Dil Factor: 1.00000

Integrator: HP RTE

Compound Sublist: 4-8260+IX.sub

Target Version: 4.04

Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

| Name | Value | Description |
|------|-------|-----------------|
| DF | 1.000 | Dilution Factor |
| Vo | 5.000 | Sample volume |

| Compounds | QUANT SIG | CONCENTRATIONS | | | | | | |
|-----------|---------------------------|----------------|-------|------------------------|---------|----------|--------|---------|
| | | MASS | RT | EXP RT | REL RT | RESPONSE | (ng) | (ug/L) |
| * | 1 Fluorobenzene | 96 | 5.041 | 5.041 (1.000) | 1815827 | 50.0000 | | |
| * | 2 Chlorobenzene-d5 | 117 | 7.680 | 7.680 (1.000) | 1616266 | 50.0000 | | |
| * | 3 1,4-Dichlorobenzene-d4 | 152 | 9.904 | 9.904 (1.000) | 657391 | 50.0000 | | |
| \$ | 4 Dibromofluoromethane | 113 | 4.485 | 4.485 (0.890) | 485939 | 57.4784 | 11.496 | |
| \$ | 5 1,2-Dichloroethane-d4 | 65 | 4.757 | 4.757 (0.944) | 662301 | 56.9707 | 11.354 | |
| \$ | 6 Toluene-d8 | 98 | 6.378 | 6.378 (0.831) | 1712090 | 44.0929 | 8.818 | |
| \$ | 7 Bromofluorobenzene | 95 | 8.780 | 8.780 (1.143) | 646832 | 39.2063 | 7.841 | |
| \$ | 8 Dichlorodifluoromethane | 85 | | Compound Not Detected. | | | | |
| \$ | 9 Chloromethane | 50 | 1.669 | 1.704 (0.331) | 17401 | 1.02663 | 0.2053 | |
| 10 | Vinyl Chloride | 62 | | Compound Not Detected. | | | | |
| 11 | Bromomethane | 94 | | Compound Not Detected. | | | | |
| 12 | Chloroethane | 64 | | Compound Not Detected. | | | | |
| 13 | Trichlorofluoromethane | 101 | | Compound Not Detected. | | | | |
| 15 | Acrolein | 56 | | Compound Not Detected. | | | | |
| 16 | Acetone | 43 | | Compound Not Detected. | | | | |
| 17 | 1,1-Dichloroethene | 96 | | Compound Not Detected. | | | | |
| 18 | Freon-113 | 151 | | Compound Not Detected. | | | | |

Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J41001A.b\UXJ24281.D
 Report Date: 04-Oct-2004 09:56

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|---------------------------------|-----------|------|----------------|--------|--------|------------------------|-------------------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 19 Iodomethane | | 142 | | | | Compound Not Detected. | |
| 20 Carbon Disulfide | | 76 | | | | Compound Not Detected. | |
| 21 Methylene Chloride | | 84 | | | | Compound Not Detected. | |
| 22 Acetonitrile | | 41 | | | | Compound Not Detected. | |
| 23 Acrylonitrile | | 53 | | | | Compound Not Detected. | |
| 24 Methyl tert-butyl ether | | 73 | | | | Compound Not Detected. | |
| 25 trans-1,2-Dichloroethene | | 96 | | | | Compound Not Detected. | |
| 26 Hexane | | 86 | | | | Compound Not Detected. | |
| 27 Vinyl acetate | | 43 | | | | Compound Not Detected. | |
| 28 1,1-Dichloroethane | | 63 | | | | Compound Not Detected. | |
| 29 tert-Butyl Alcohol | | 59 | | | | Compound Not Detected. | |
| 30 2-Butanone | | 43 | | | | Compound Not Detected. | |
| M 31 1,2-Dichloroethane (total) | | 96 | | | | Compound Not Detected. | |
| 32 cis-1,2-dichloroethene | | 96 | | | | Compound Not Detected. | |
| 33 2,2-Dichloropropane | | 77 | | | | Compound Not Detected. | |
| 34 Bromochloromethane | | 128 | | | | Compound Not Detected. | |
| 35 Chloroform | | 83 | | | | Compound Not Detected. | |
| 36 Tetrahydrofuran | | 42 | | | | Compound Not Detected. | |
| 37 1,1,1-Trichloroethane | | 97 | | | | Compound Not Detected. | |
| 38 1,1-Dichloropropene | | 75 | | | | Compound Not Detected. | |
| 39 Carbon Tetrachloride | | 117 | | | | Compound Not Detected. | |
| 40 1,2-Dichloroethane | | 62 | | | | Compound Not Detected. | |
| 41 Benzene | | 78 | | | | Compound Not Detected. | |
| 42 Trichloroethene | | 130 | | | | Compound Not Detected. | |
| 43 1,2-Dichloropropane | | 63 | | | | Compound Not Detected. | |
| 44 1,4-Dioxane | | 88 | | | | Compound Not Detected. | |
| 45 Dibromomethane | | 93 | | | | Compound Not Detected. | |
| 46 Bromodichloromethane | | 83 | | | | Compound Not Detected. | |
| 47 2-Chloroethyl vinyl ether | | 63 | | | | Compound Not Detected. | |
| 48 cis-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | |
| 49 4-Methyl-2-pentanone | | 43 | | | | Compound Not Detected. | |
| 50 Toluene | | 91 | | | | Compound Not Detected. | |
| 51 trans-1,3-Dichloropropene | | 75 | | | | Compound Not Detected. | |
| 52 Ethyl Methacrylate | | 69 | | | | Compound Not Detected. | |
| 53 1,1,2-Trichloroethane | | 97 | | | | Compound Not Detected. | |
| 54 1,3-Dichloropropane | | 76 | | | | Compound Not Detected. | |
| 55 Tetrachloroethene | | 164 | | | | Compound Not Detected. | |
| 56 2-Hexanone | | 43 | | | | Compound Not Detected. | |
| 57 Dibromochloromethane | | 129 | | | | Compound Not Detected. | |
| 58 1,2-Dibromoethane | | 107 | | | | Compound Not Detected. | |
| 59 Chlorobenzene | | 112 | | | | Compound Not Detected. | |
| 60 1,1,1,2-Tetrachloroethane | | 131 | | | | Compound Not Detected. | |
| 61 Ethylbenzene | | 106 | | | | Compound Not Detected. | |
| 62 m + p-Xylene | | 106 | | | | Compound Not Detected. | |
| M 63 Xylenes (total) | | 106 | | | | Compound Not Detected. | |
| 64 Xylene-o | | 106 | | | | Compound Not Detected. | |
| 65 Styrene | | 104 | | | | Compound Not Detected. | |

Data File: \\qcanch04\dd\chem\MSV\ a3ux11.i \J41001A.b\UXJ24281.D
 Report Date: 04-Oct-2004 09:56

| Compounds | QUANT SIG | MASS | CONCENTRATIONS | | | | |
|--------------------------------|-----------|------|----------------|--------|--------|------------------------|-------------------------------|
| | | | RT | EXP RT | REL RT | RESPONSE | ON-COLUMN (ng) FINAL (ug/L) |
| 66 Bromoform | | 173 | | | | Compound Not Detected. | |
| 67 Isopropylbenzene | | 105 | | | | Compound Not Detected. | |
| 68 1,1,2,2-Tetrachloroethane | | 83 | | | | Compound Not Detected. | |
| 69 1,4-Dichloro-2-butene | | 53 | | | | Compound Not Detected. | |
| 70 1,2,3-Trichloropropane | | 110 | | | | Compound Not Detected. | |
| 71 Bromobenzene | | 156 | | | | Compound Not Detected. | |
| 72 n-Propylbenzene | | 120 | | | | Compound Not Detected. | |
| 73 2-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 74 1,3,5-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 75 4-Chlorotoluene | | 126 | | | | Compound Not Detected. | |
| 76 tert-Butylbenzene | | 119 | | | | Compound Not Detected. | |
| 77 1,2,4-Trimethylbenzene | | 105 | | | | Compound Not Detected. | |
| 78 sec-Butylbenzene | | 105 | | | | Compound Not Detected. | |
| 79 4-Isopropyltoluene | | 119 | | | | Compound Not Detected. | |
| 80 1,3-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 81 1,4-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 82 n-Butylbenzene | | 91 | | | | Compound Not Detected. | |
| 83 1,2-Dichlorobenzene | | 146 | | | | Compound Not Detected. | |
| 84 1,2-Dibromo-3-chloropropane | | 157 | | | | Compound Not Detected. | |
| 85 1,2,4-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 86 Hexachlorobutadiene | | 225 | | | | Compound Not Detected. | |
| 87 Naphthalene | | 128 | | | | Compound Not Detected. | |
| 88 1,2,3-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |
| 14 Dichlorofluoromethane | | 67 | | | | Compound Not Detected. | |
| 89 Ethyl Ether | | 59 | | | | Compound Not Detected. | |
| 91 3-Chloropropene | | 76 | | | | Compound Not Detected. | |
| 92 Isopropyl Ether | | 87 | | | | Compound Not Detected. | |
| 93 2-Chloro-1,3-butadiene | | 53 | | | | Compound Not Detected. | |
| 94 Propionitrile | | 54 | | | | Compound Not Detected. | |
| 95 Ethyl Acetate | | 43 | | | | Compound Not Detected. | |
| 96 Methacrylonitrile | | 41 | | | | Compound Not Detected. | |
| 97 Isobutanol | | 41 | | | | Compound Not Detected. | |
| 99 n-Butanol | | 56 | | | | Compound Not Detected. | |
| 100 Methyl Methacrylate | | 41 | | | | Compound Not Detected. | |
| 101 2-Nitropropane | | 41 | | | | Compound Not Detected. | |
| 103 Cyclohexanone | | 55 | | | | Compound Not Detected. | |
| 98 Cyclohexane | | 56 | | | | Compound Not Detected. | |
| 143 Methyl Acetate | | 43 | | | | Compound Not Detected. | |
| 144 Methylcyclohexane | | 83 | | | | Compound Not Detected. | |
| 141 1,3,5-Trichlorobenzene | | 180 | | | | Compound Not Detected. | |

Data File: \\pcanch04\dd\chem\MSV\z3ux11.1\J41001A.b\UXJ24281.D

Date : 01-OCT-2004 09:32

Client ID:

Instrument: z3ux11.i

Sample Info: VELK

Operator: 43582

Purge Volume: 5.0

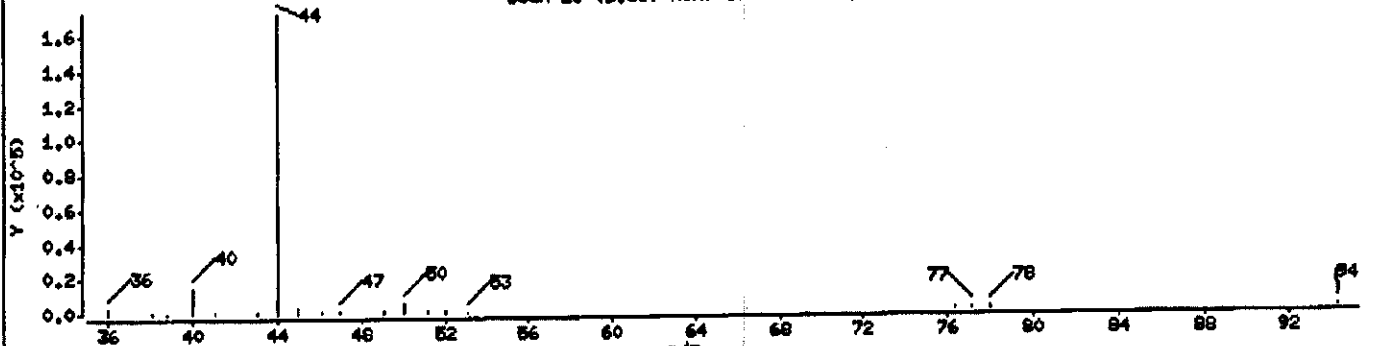
Column diameter: 0.18

Column phase: DB624

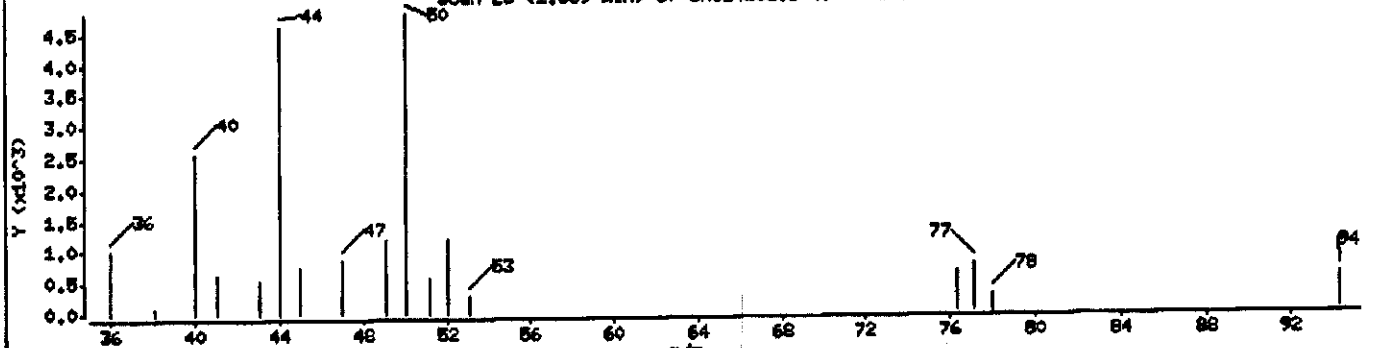
Concentration: 0.2053 ug/L

9 Chloromethane

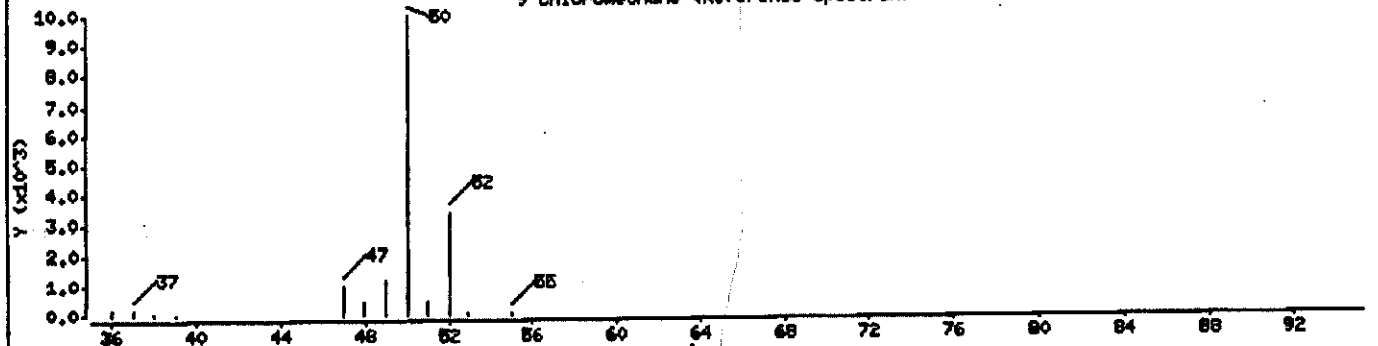
Scan 25 (1.669 min) of UXJ24281.D



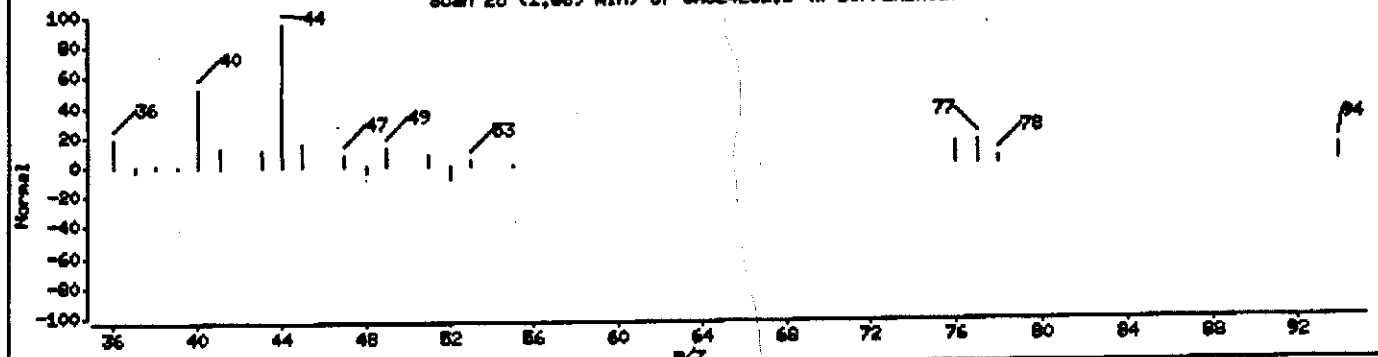
Scan 25 (1.669 min) of UXJ24281.D (Subtracted)



9 Chloromethane (Reference Spectrum)



Scan 25 (1.669 min) of UXJ24281.D (% DIFFERENCE)



Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J41001A.b\UXJ24281.D
Report Date: 04-Oct-2004 09:56

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J41001A.b\UXJ24281.D

Lab Smp Id: grkvplaa

Inj Date : 01-OCT-2004 09:32

Inst ID: A3UX11.i

Operator : 43582

Smp Info : VBLK

Misc Info : J41001A,8260LLUX11,,43582,3,,BLANK,,0

Comment :

Method : \\QCANOHO4\dd\chem\MSV\A3UX11.i\J41001A.b\8260LLUX11.m

Meth Date : 04-Oct-2004 09:54 evans1 Quant Type: ISTD

Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D

Als bottle: 5 QC Sample: BLANK

Dil Factor: 1.00000

Integrator: HP RTE

Compound Sublist: 4-8260+IX.sub

Target Version: 4.04

Processing Host: CANPMSV07

- NO TENTATIVELY IDENTIFIED COMPOUNDS -

SEVERN
TRENT

STL

MISCELLANEOUS DATA

UX11
Batch # _____

**STL-North Canton
GC/MS VOA Run Log**

Date: 8/16/07

| Column | | BFB | Analysis | | Purge & Trap |
|---------------------|----------------------|---------------------|---------------------|-----------------------|--------------------------|
| Type: DB624 | | 100 C for 0.1 min | 45 C for 2 min | Trap: #10 | |
| Length 20 M | | to 200 C @ 20 C/min | to 200 C @ 20 C/min | Purge: 11 | |
| I.D. 0.18 mm | | Hold 0 min | Hold 0 min | Desorb: 1 min @ 240 C | |
| Flow Rate 0.4ml/min | | IS # 12149 SS # | Hold 3 min | Bake: 5 min @ 250 C | |
| Auto num | Sample ID Workorder# | Method | File Name | Amt purged | Std number / Sample prep |
| | | BFB | BFB 207 | SONG | dinnyret (13/09) |
| | 101 STD | | WTJ3202 | 200mg | 12149, 50, 53, 60 |
| | 101 STD | | | 03 | 100 mg |
| | 101 STD | | | 04 | SONG |
| | 101 STD | | | 05 | 25mg |
| | 101 STD | | | 06 | long |
| | 101 STD | | | 07 | SONG |
| | ACV | | | 08 | SONG values |
| | AG STD | | | 09 | 200mg |
| | AG STD | | | 10 | 100 mg |
| | AG STD | | | 11 | SONG |
| | AG STD | | | 12 | 25mg |
| | AG STD | | | 13 | long |
| | AG STD | | | 14 | SONG |

Analyst: JR
Level 2 review: JR

UX11

Batch # 4259349

STL-North Canton
GC/MS VOA Run Log

(9/16) Date: 9/14/04

Column
 Type: DB624
 Length 20 M
 I.D. 0.18 mm
 Flow Rate 0.4ml/min

BFB
 100 C for 0.1 min
 to 200 C @ 20 C/min
 Hold 2 min

IS # 12246 ss # 12247

Analysis
 45 C for 2 min
 to 200 C @ 20 C/min
 to 250 C @ 20 C/min
 Hold 3 min

Purge & Trap
 Trap: #10
 Purge: 11
 Desorb: 1 min @ 240 C
 Bake: 5 min @ 250 C
 Heated purge: Yes No

| Auto num | Sample ID / Workorder# | Method | File Name | Amt purged | Std number / Sample prep | Comments | Sample status |
|----------|------------------------|--------|-----------|------------|--------------------------|--------------|---------------|
| ✓ | BRB | | DPB 232 | 50mg | +50ng | (13:21) | On |
| ✓ | 101 STD | | UKJ23870 | 200mg | V2252,66 | | On |
| ✓ | 101 STD | | 71 | 100mg | 53,55 | | On |
| ✓ | 101 STD | - | 72 | 50mg | | J40914 | On |
| ✓ | 101 STD | | 73 | 25mg | | | On |
| ✓ | 101 STD | | 74 | 10 mg | | | On |
| ✓ | 101 STD | | 75 | 5 mg | | | On |
| ✓ | PCV | | 76 | 50mg | V2259 | | On |
| ✓ | Check GP810W | | 77 | 1 | | | On |
| ✓ | Check Dsp | | 78 | 1 | | | On |
| ✓ | Blank | | 79 | Sme | | | On |
| ✓ | GPMLIAA | | 80 | 0.2mg/ml | | | On |
| ✓ | GPMLIAA (S) | | 81 | | +50ng | | On |
| ✓ | GPMLIAA (O) | | 82 | | 1 | | On |
| ✓ | GPMLIAA ED | | 83 | Sme | | | On |
| ✓ | GPHTDIAZ | | 84 | | | | On |
| ✓ | GPJWVIAA ED | | 85 | | | | On |
| ✓ | GPJW11AA | | 86 | | | | On |
| ✓ | GPJW31AA | | 87 | | | | On |
| ✓ | GPJW41AA | | 88 | | | | On |
| ✓ | GPJW51AA | | 89 | | | | On |
| ✓ | GPJW61AA | | 90 | | | | On |
| ✓ | GPJW71AA | | 91 | | | | On |
| ✓ | GPJW81AA | | 92 | | | | On |
| ✓ | GPJW91AA | | 93 | | | | On |
| ✓ | GPKLEIAA | 6m | 84 | | | | On |
| ✓ | GPPLFIAA | | 94 | | | | On |
| ✓ | GPKMJIAA | | 95 | | | R 1.5ml vial | OK |
| ✓ | GPLODJIAA | m | 96 | | | | On |
| ✓ | GPLDPAA | m | 97 | | | | OK |
| ✓ | GPLOTIAA | m | 98 | | | | OK |
| ✓ | GPLOTTIAA | b | 99 | | | | OK |

Analyst: *[Signature]*
 Level 2 review: *[Signature]*

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UX11
Batch # 4275213STL-North Canton
GC/MS VOA Run LogDate: 10/1/04

Column
Type: DB624
Length: 20 M
I.D.: 0.18 mm
Flow Rate: 0.4ml/min

BFB
100 C for 0.1 min
to 200 C @ 20 C/min
Hold = min

IS# V2313 ss# V2314

Analysis
45 C for 2 min
to 200 C @ 20 C/min
to = C @ = C/min
Hold 3 min

Purge & Trap
Trap: #10
Purge: 11
Desorb: 1 min @ 240 C
Bake: 5 min @ 250 C
Heated purge: Yes No

| Sample ID | Sample Name | Conc | Notes | Run |
|------------------|-------------|------|----------------------|--------------------|
| 1 BPB | BPB246 | 50ug | dir inj ext | (9:30) |
| 1-1 W1 STD | WT24697 | | V2313, 09/08/04 0974 | 09 |
| 1-2 A2 STD | 78 | | V2313 J4081b | 09 |
| 1-3 Check | GRKVP | 79 | M329 | 09 |
| 1-4 Check Dsp | | 80 | | 09 |
| 1-5 blank | | 81 | SMN | 09 |
| 1-6 GQ2KWAIA | | 82 | 0.005mg/ml | OK @ 50% |
| 1-7 GQ2LVIAA | | 83 | 0.005mg/ml | 2AA @ 50% |
| 1-8 GQ2UNIAA | | 84 | 0.005mg/ml | 2AA @ 50% |
| 1-9 GQ2LFIAA | | 85 | 0.5mg/ml | foams! (no 10x) ON |
| 1-10 GROVE1AA | Rush | 86 | SMN | ON |
| 1-11 GROXI1AA | | 87 | | ON |
| 1-12 GROXG1AA | | 88 | | ON |
| 1-13 GROOD1AA | | 89 | | ON |
| 1-14 GQ2L21AA | | 90 | SMN | ON |
| 1-15 GQ2UF1AA | | 91 | | ON |
| 1-16 GQ2LH1AA | | 92 | | ON |
| 1-17 GQ2LK1AA | | 93 | | ON |
| 1-18 GQ2LC1AA | | 94 | | ON |
| 1-19 GQ2LQ1AA | | 95 | | ON |
| 1-20 GQ2LI1AA | | 96 | | ON |
| 1-21 GQ2LY1AA | | 97 | | ON |
| 1-22 GQ2LS1AA | | 98 | | ON |
| 1-23 GQ2KL61A(S) | | 99 | +30% | ON |
| 1-24 GQ2KL61A(D) | | 100 | | ON |
| 1-25 GQ2LG1AA | | 01 | SMN | ON |
| 1-26 GQ2L91AA | | 02 | | ON |
| 1-27 GQ2KL61AA | | 03 | SMN | ON |
| 1-28 GQ2KL7AA | | 04 | 0.07mg/ml | ON |

10/4/04Analyst: J. Smith
Level 2 review: SPM

MSVOC

Lot Summary - A4I290193

SDG:

(4129) (93) (53)

CLIENT: 5670 PAYNE FIRM INC.

PROJECT MANAGER: Roger K. Toth

SITE: EM SCIENCE (OH)

LOT COMMENTS:

QC PACKAGE: Expanded Deliverables

R E P R I N T - - - - -

= Field(s) Changed

| SAMP# | W/O NO. | PARAMETER | X-REF | Sampled | Expires | Est | Sample ID, Comments / Analysis | Comments |
|-------|---------|-----------|-------|---------|---------|-----|--------------------------------|----------|
|-------|---------|-----------|-------|---------|---------|-----|--------------------------------|----------|

| | | | | | |
|------|----------------------------------|---------|----------|---|---|
| 001- | GRDVE-1AA XX I 25 QK 01 MS8260LL | 9/28/04 | 10/12/04 | Y | #VE540/21-26/092804 Q: CLP MSVOA TCL Standard List #EXP DEL, SDG #4I29193 (CL), RUSH DUE FRI 10-1-04, VOC #NEEDS 10X LESSER DILUTION. AP9 Compounds |
| 002- | GRDX1-1AA XX I 25 QK 01 MS8260LL | 9/28/04 | 10/12/04 | Y | #VE540/35.5-40.5/092804 Q: CLP MSVOA TCL Standard List #EXP DEL, SDG #4I29193 (CL), RUSH DUE FRI 10-1-04, VOC #NEEDS 10X LESSER DILUTION. AP9 Compounds |
| 003- | GRDX6-1AA XX I 25 QK 01 MS8260LL | 9/28/04 | 10/12/04 | Y | #VE541/26.5-31.5/092804 Q: CLP MSVOA TCL Standard List #EXP DEL, SDG #4I29193 (CL), RUSH DUE FRI 10-1-04, VOC #NEEDS 10X LESSER DILUTION. AP9 Compounds |
| 004- | GRDOD-1AA XX I 25 QK 01 MS8260LL | 9/28/04 | 10/12/04 | Y | #TRIP BLANK/092804 Q: CLP MSVOA TCL Standard List #EXP DEL, SDG #4I29193 (CL), RUSH DUE FRI 10-1-04, VOC #NEEDS 10X LESSER DILUTION. AP9 Compounds |

| LOT NUMBER | LAB SAMPLE ID | ANALYSIS TYPE | ANALYSIS DATE | ANALYST |
|---------------|---------------------|------------------|------------------|-------------|
| A41290193 | 1 GRDURIAA | MS8260LL | 10/01/04 | Laura Evans |
| A41290193 | 2 GRDXK1MA | MS8260LL | 10/01/04 | Laura Evans |
| A41290193 | 3 GRDXG1MA | MS8260LL | 10/01/04 | Laura Evans |
| A41290193 | 4 GRDD01AA | MS8260LL | 10/01/04 | Laura Evans |

* * * E N D O F R E P O R T * * *

END OF REPORT